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(54) Title: METHODS FOR GENERATING POLYNUCLEOTIDES HAVING DESIRED CHARACTERISTICS BY ITERATIVE
SELECTION AND RECOMBINATION

(57) Abstract

A method for DNA reassembly after random fragmentation, and its application to mutagenesis of nucleic acid sequences by *in vitro* or *in vivo* recombination is described. In particular, a method for the production of nucleic acid fragments or polynucleotides encoding mutant proteins is described. The present invention also relates to a method of repeated cycles of mutagenesis, shuffling and selection which allow for the directed molecular evolution *in vitro* or *in vivo* of proteins.

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5 METHODS FOR GENERATING POLYNUCLEOTIDES HAVING DESIRED
CHARACTERISTICS BY ITERATIVE SELECTION AND RECOMBINATION

Cross-Reference to Related Applications

The present application is a continuation-in-part of USSN 08/621,859, filed March 25, 1996, and a continuation-in-part of 10 USSN 08/564,955, filed November 30, 1995.

Field of the Invention

The present invention relates to a method for the production of polynucleotides conferring a desired phenotype and/or encoding 15 a protein having an advantageous predetermined property which is selectable or can be screened for. In an aspect, the method is used for generating and selecting or screening for desired nucleic acid fragments encoding mutant proteins.

20 BACKGROUND AND DESCRIPTION OF RELATED ART

The complexity of an active sequence of a biological macromolecule, e.g. proteins, DNA etc., has been called its information content ("IC"; 5-9). The information content of a protein has been defined as the resistance of the active protein 25 to amino acid sequence variation, calculated from the minimum number of invariable amino acids (bits) required to describe a family of related sequences with the same function (9, 10). Proteins that are sensitive to random mutagenesis have a high information content. In 1974, when this definition was coined, 30 protein diversity existed only as taxonomic diversity.

Molecular biology developments such as molecular libraries have allowed the identification of a much larger number of variable bases, and even to select functional sequences from random libraries. Most residues can be varied, although 35 typically not all at the same time, depending on compensating changes in the context. Thus a 100 amino acid protein can contain only 2,000 different mutations, but 20^{100} possible combinations of mutations.

Information density is the Information Content/unit length of a sequence. Active sites of enzymes tend to have a high information density. By contrast, flexible linkers in enzymes have a low information density (8).

- 5 Current methods in widespread use for creating mutant proteins in a library format are error-prone polymerase chain reaction (11, 12, 19) and cassette mutagenesis (8, 20, 21, 22, 40, 41, 42), in which the specific region to be optimized is replaced with a synthetically mutagenized oligonucleotide.
- 10 Alternatively, mutator strains of host cells have been employed to add mutational frequency (Greener and Callahan (1995) Strategies in Mol. Biol. 7: 32). In each case, a 'mutant cloud' (4) is generated around certain sites in the original sequence.

Error-prone PCR uses low-fidelity polymerization conditions to introduce a low level of point mutations randomly over a long sequence. Error prone PCR can be used to mutagenize a mixture of fragments of unknown sequence. However, computer simulations have suggested that point mutagenesis alone may often be too gradual to allow the block changes that are required for continued sequence evolution. The published error-prone PCR protocols are generally unsuited for reliable amplification of DNA fragments greater than 0.5 to 1.0 kb, limiting their practical application. Further, repeated cycles of error-prone PCR lead to an accumulation of neutral mutations, which, for example, may make a protein immunogenic.

In oligonucleotide-directed mutagenesis, a short sequence is replaced with a synthetically mutagenized oligonucleotide. This approach does not generate combinations of distant mutations and is thus not significantly combinatorial.

30 The limited library size relative to the vast sequence length means that many rounds of selection are unavoidable for protein optimization. Mutagenesis with synthetic oligonucleotides requires sequencing of individual clones after each selection round followed by grouping into families, arbitrarily choosing

35 a single family, and reducing it to a consensus motif, which is resynthesized and reinserted into a single gene followed by additional selection. This process constitutes a statistical

bottleneck, it is labor intensive and not practical for many rounds of mutagenesis.

Error-prone PCR and oligonucleotide-directed mutagenesis are thus useful for single cycles of sequence fine tuning but rapidly become limiting when applied for multiple cycles.

Error-prone PCR can be used to mutagenize a mixture of fragments of unknown sequence (11, 12). However, the published error-prone PCR protocols (11, 12) suffer from a low processivity of the polymerase. Therefore, the protocol is very difficult to employ for the random mutagenesis of an average-sized gene. This inability limits the practical application of error-prone PCR.

Another serious limitation of error-prone PCR is that the rate of down-mutations grows with the information content of the sequence. At a certain information content, library size, and mutagenesis rate, the balance of down-mutations to up-mutations will statistically prevent the selection of further improvements (statistical ceiling).

Finally, repeated cycles of error-prone PCR will also lead to the accumulation of neutral mutations, which can affect, for example, immunogenicity but not binding affinity.

Thus error-prone PCR was found to be too gradual to allow the block changes that are required for continued sequence evolution (1, 2).

In cassette mutagenesis, a sequence block of a single template is typically replaced by a (partially) randomized sequence. Therefore, the maximum information content that can be obtained is statistically limited by the number of random sequences (i.e., library size). This constitutes a statistical bottleneck, eliminating other sequence families which are not currently best, but which may have greater long term potential.

Further, mutagenesis with synthetic oligonucleotides requires sequencing of individual clones after each selection round (20). Therefore, this approach is tedious and is not practical for many rounds of mutagenesis.

Error-prone PCR and cassette mutagenesis are thus best suited and have been widely used for fine-tuning areas of

comparatively low information content. An example is the selection of an RNA ligase ribozyme from a random library using many rounds of amplification by error-prone PCR and selection (13).

5 It is becoming increasingly clear our scientific tools for the design of recombinant linear biological sequences such as protein, RNA and DNA are not suitable for generating the necessary sequence diversity needed to optimize many desired properties of a macromolecule or organism. Finding better and
10 better mutants depends on searching more and more sequences within larger and larger libraries, and increasing numbers of cycles of mutagenic amplification and selection are necessary. However as discussed above, the existing mutagenesis methods that
15 are in widespread use have distinct limitations when used for repeated cycles.

Evolution of most organisms occurs by natural selection and sexual reproduction. Sexual reproduction ensures mixing and combining of the genes of the offspring of the selected individuals. During meiosis, homologous chromosomes from the
20 parents line up with one another and cross-over part way along their length, thus swapping genetic material. Such swapping or shuffling of the DNA allows organisms to evolve more rapidly (1, 2). In sexual recombination, because the inserted sequences were of proven utility in a homologous environment, the inserted
25 sequences are likely to still have substantial information content once they are inserted into the new sequence.

Marton et al., (27) describes the use of PCR *in vitro* to monitor recombination in a plasmid having directly repeated sequences. Marton et al. discloses that recombination will occur
30 during PCR as a result of breaking or nicking of the DNA. This will give rise to recombinant molecules. Meyerhans et al. (23) also disclose the existence of DNA recombination during *in vitro* PCR.

The term Applied Molecular Evolution ("AME") means the
35 application of an evolutionary design algorithm to a specific, useful goal. While many different library formats for AME have been reported for polynucleotides (3, 11-14), peptides and

proteins (phage (15-17), lacI (18) and polysomes, in none of these formats has recombination by random cross-overs been used to deliberately create a combinatorial library.

Theoretically there are 2,000 different single mutants 5 of a 100 amino acid protein. A protein of 100 amino acids has 20^{100} possible combinations of mutations, a number which is too large to exhaustively explore by conventional methods. It would be advantageous to develop a system which would allow the generation and screening of all of these possible combination 10 mutations.

Winter and coworkers (43,44) have utilized an *in vivo* site specific recombination system to combine light chain antibody genes with heavy chain antibody genes for expression in a phage system. However, their system relies on specific sites 15 of recombination and thus is limited. Hayashi et al. (48) report simultaneous mutagenesis of antibody CDR regions in single chain antibodies (scFv) by overlap extension and PCR.

Caren et al. (45) describe a method for generating a large population of multiple mutants using random *in vivo* 20 recombination. However, their method requires the recombination of two different libraries of plasmids, each library having a different selectable marker. Thus the method is limited to a finite number of recombinations equal to the number of selectable 25 markers existing, and produces a concomitant linear increase in the number of marker genes linked to the selected sequence(s). Caren et al. does not describe the use of multiple selection cycles; recombination is used solely to construct larger libraries.

Calogero et al. (46) and Galizzi et al. (47) report 30 that *in vivo* recombination between two homologous but truncated insect-toxin genes on a plasmid can produce a hybrid gene. Radman et al. (49) report *in vivo* recombination of substantially mismatched DNA sequences in a host cell having defective mismatch repair enzymes, resulting in hybrid molecule formation.

It would be advantageous to develop a method for the 35 production of mutant proteins which method allowed for the development of large libraries of mutant nucleic acid sequences

which were easily search d. The invention described herein is directed to the use of repeated cycles of point mutagenesis, nucleic acid shuffling and selection which allow for the directed molecular evolution *in vitro* of highly complex linear sequences, 5 such as proteins through random recombination.

Accordingly, it would be advantageous to develop a method which allows for the production of large libraries of mutant DNA, RNA or proteins and the selection of particular mutants for a desired goal. The invention described herein is 10 directed to the use of repeated cycles of mutagenesis, *in vivo* recombination and selection which allow for the directed molecular evolution *in vivo* and *in vitro* of highly complex linear sequences, such as DNA, RNA or proteins through recombination.

15 Further advantages of the present invention will become apparent from the following description of the invention with reference to the attached drawings.

SUMMARY OF THE INVENTION

20 The present invention is directed to a method for generating a selected polynucleotide sequence or population of selected polynucleotide sequences, typically in the form of amplified and/or cloned polynucleotides, whereby the selected polynucleotide sequence(s) possess a desired phenotypic 25 characteristic (e.g., encode a polypeptide, promote transcription of linked polynucleotides, bind a protein, and the like) which can be selected for. One method of identifying polypeptides that possess a desired structure or functional property, such as binding to a predetermined biological macromolecule (e.g., a 30 receptor), involves the screening of a large library of polypeptides for individual library members which possess the desired structure or functional property conferred by the amino acid sequence of the polypeptide.

In a general aspect, the invention provides a method, 35 termed "sequence shuffling", for generating libraries of recombinant polynucleotides having a desired characteristic which can be selected or screened for. Libraries of recombinant

polynucleotides are generated from a population of related-sequence polynucleotides which comprise sequence regions which have substantial sequence identity and can be homologously recombined in vitro or in vivo. In the method, at least two species of the related-sequence polynucleotides are combined in a recombination system suitable for generating sequence-recombined polynucleotides, wherein said sequence-recombined polynucleotides comprise a portion of at least one first species of a related-sequence polynucleotide with at least one adjacent portion of at least one second species of a related-sequence polynucleotide. Recombination systems suitable for generating sequence-recombined polynucleotides can be either: (1) in vitro systems for homologous recombination or sequence shuffling via amplification or other formats described herein, or (2) in vivo systems for homologous recombination or site-specific recombination as described herein. The population of sequence-recombined polynucleotides comprises a subpopulation of polynucleotides which possess desired or advantageous characteristics and which can be selected by a suitable selection or screening method. The selected sequence-recombined polynucleotides, which are typically related-sequence polynucleotides, can then be subjected to at least one recursive cycle wherein at least one selected sequence-recombined polynucleotide is combined with at least one distinct species of related-sequence polynucleotide (which may itself be a selected sequence-recombined polynucleotide) in a recombination system suitable for generating sequence-recombined polynucleotides, such that additional generations of sequence-recombined polynucleotide sequences are generated from the selected sequence-recombined polynucleotides obtained by the selection or screening method employed. In this manner, recursive sequence recombination generates library members which are sequence-recombined polynucleotides possessing desired characteristics. Such characteristics can be any property or attribute capable of being selected for or detected in a screening system, and may include properties of: an encoded protein, a transcriptional element, a sequence controlling transcription, RNA processing, RNA

stability, chromatin conformation, translation, or other expression property of a gene or transgene, a replicative element, a protein-binding element, or the like, such as any feature which confers a selectable or detectable property.

The present invention provides a method for generating libraries of displayed polypeptides or displayed antibodies suitable for affinity interaction screening or phenotypic screening. The method comprises (1) obtaining a first plurality of selected library members comprising a displayed polypeptide or displayed antibody and an associated polynucleotide encoding said displayed polypeptide or displayed antibody, and obtaining said associated polynucleotides or copies thereof wherein said associated polynucleotides comprise a region of substantially identical sequence, optionally introducing mutations into said polynucleotides or copies, and (2) pooling and fragmenting, by nuclease digestion, partial extension PCR amplification, PCR stuttering, or other suitable fragmenting means, typically producing random fragments or fragment equivalents, said associated polynucleotides or copies to form fragments thereof under conditions suitable for PCR amplification, performing PCR amplification and optionally mutagenesis, and thereby homologously recombining said fragments to form a shuffled pool of recombined polynucleotides, whereby a substantial fraction (e.g., greater than 10 percent) of the recombined polynucleotides of said shuffled pool are not present in the first plurality of selected library members, said shuffled pool composing a library of displayed polypeptides or displayed antibodies suitable for affinity interaction screening. Optionally, the method comprises the additional step of screening the library members of the shuffled pool to identify individual shuffled library members having the ability to bind or otherwise interact (e.g., such as catalytic antibodies) with a predetermined macromolecule, such as for example a proteinaceous receptor, peptide, oligosaccharide, virion, or other predetermined compound or structure. The displayed polypeptides, antibodies, peptidomimetic antibodies, and variable region sequences that are identified from such libraries can be used for therapeutic, diagnostic,

research, and related purposes (e.g., catalysts, solutes for increasing osmolarity of an aqueous solution, and the like), and/or can be subjected to one or more additional cycles of shuffling and/or affinity selection. The method can be modified 5 such that the step of selecting is for a phenotypic characteristic other than binding affinity for a predetermined molecule (e.g., for catalytic activity, stability, oxidation resistance, drug resistance, or detectable phenotype conferred on a host cell).

10 In one embodiment, the first plurality of selected library members is fragmented and homologously recombined by PCR in vitro. Fragment generation is by nuclease digestion, partial extension PCR amplification, PCR stuttering, or other suitable fragmenting means, such as described herein. Stuttering is 15 fragmentation by incomplete polymerase extension of templates. A recombination format based on very short PCR extension times was employed to create partial PCR products, which continue to extend off a different template in the next (and subsequent) cycle(s).

20 In one embodiment, the first plurality of selected library members is fragmented in vitro, the resultant fragments transferred into a host cell or organism and homologously recombined to form shuffled library members in vivo.

25 In one embodiment, the first plurality of selected library members is cloned or amplified on episomally replicable vectors, a multiplicity of said vectors is transferred into a cell and homologously recombined to form shuffled library members in vivo.

30 In one embodiment, the first plurality of selected library members is not fragmented, but is cloned or amplified on an episomally replicable vector as a direct repeat or indirect (or inverted) repeat, which each repeat comprising a distinct species of selected library member sequence, said vector is transferred into a cell and homologously recombined by intra- 35 vector or inter-vector recombination to form shuffled library members in vivo.

In an embodiment, combinations of in vitro and in vivo shuffling are provided to enhance combinatorial diversity.

The present invention provides a method for generating libraries of displayed antibodies suitable for affinity interaction screening. The method comprises (1) obtaining a first plurality of selected library members comprising a displayed antibody and an associated polynucleotide encoding said displayed antibody, and obtaining said associated polynucleotides or copies thereof, wherein said associated polynucleotides 5 comprise a region of substantially identical variable region framework sequence, and (2) pooling and fragmenting said associated polynucleotides or copies to form fragments thereof under conditions suitable for PCR amplification and thereby 10 homologously recombining said fragments to form a shuffled pool of recombined polynucleotides comprising novel combinations of CDRs, whereby a substantial fraction (e.g., greater than 10 percent) of the recombined polynucleotides of said shuffled pool 15 comprise CDR combinations which are not present in the first plurality of selected library members, said shuffled pool composing a library of displayed antibodies comprising CDR permutations and suitable for affinity interaction screening. 20 Optionally, the shuffled pool is subjected to affinity screening to select shuffled library members which bind to a predetermined epitope (antigen) and thereby selecting a plurality of selected 25 shuffled library members. Optionally, the plurality of selected shuffled library members can be shuffled and screened iteratively, from 1 to about 1000 cycles or as desired until library members having a desired binding affinity are obtained.

Accordingly, one aspect of the present invention 30 provides a method for introducing one or more mutations into a template double-stranded polynucleotide, wherein the template double-stranded polynucleotide has been cleaved or PCR amplified (via partial extension or stuttering) into random fragments of a desired size, by adding to the resultant population of double- 35 stranded fragments one or more single or double-stranded oligonucleotides, wherein said oligonucleotides comprise an area of identity and an area of heterology to the template

polynucleotide; denaturing the resultant mixture of double-stranded random fragments and oligonucleotides into single-stranded fragments; incubating the resultant population of single-stranded fragments with a polymerase under conditions which result in the annealing of said single-stranded fragments at regions of identity between the single-stranded fragments and formation of a mutagenized double-stranded polynucleotide; and repeating the above steps as desired.

In another aspect the present invention is directed to a method of producing recombinant proteins having biological activity by treating a sample comprising double-stranded template polynucleotides encoding a wild-type protein under conditions which provide for the cleavage of said template polynucleotides into random double-stranded fragments having a desired size; adding to the resultant population of random fragments one or more single or double-stranded oligonucleotides, wherein said oligonucleotides comprise areas of identity and areas of heterology to the template polynucleotide; denaturing the resultant mixture of double-stranded fragments and oligonucleotides into single-stranded fragments; incubating the resultant population of single-stranded fragments with a polymerase under conditions which result in the annealing of said single-stranded fragments at the areas of identity and formation of a mutagenized double-stranded polynucleotide; repeating the above steps as desired; and then expressing the recombinant protein from the mutagenized double-stranded polynucleotide.

A third aspect of the present invention is directed to a method for obtaining a chimeric polynucleotide by treating a sample comprising different double-stranded template polynucleotides wherein said different template polynucleotides contain areas of identity and areas of heterology under conditions which provide for the cleavage of said template polynucleotides into random double-stranded fragments of a desired size; denaturing the resultant random double-stranded fragments contained in the treated sample into single-stranded fragments; incubating the resultant single-strand d fragments with polymerase under conditions which provide for th annealing

of the single-stranded fragments at the areas of identity and the formation of a chimeric double-stranded polynucleotide sequence comprising template polynucleotide sequences; and repeating the above steps as desired.

5 A fourth aspect of the present invention is directed to a method of replicating a template polynucleotide by combining *in vitro* single-stranded template polynucleotides with small random single-stranded fragments resulting from the cleavage and denaturation of the template polynucleotide, and incubating said
10 mixture of nucleic acid fragments in the presence of a nucleic acid polymerase under conditions wherein a population of double-stranded template polynucleotides is formed.

The invention also provides the use of polynucleotide shuffling, *in vitro* and/or *in vivo* to shuffle polynucleotides
15 encoding polypeptides and/or polynucleotides comprising transcriptional regulatory sequences.

The invention also provides the use of polynucleotide shuffling to shuffle a population of viral genes (e.g., capsid proteins, spike glycoproteins, polymerases, proteases, etc.) or
20 viral genomes (e.g., paramyxoviridae, orthomyxoviridae, herpesviruses, retroviruses, reoviruses, rhinoviruses, etc.). In an embodiment, the invention provides a method for shuffling sequences encoding all or portions of immunogenic viral proteins
25 to generate novel combinations of epitopes as well as novel epitopes created by recombination; such shuffled viral proteins may comprise epitopes or combinations of epitopes which are likely to arise in the natural environment as a consequence of viral evolution (e.g., such as recombination of influenza virus strains).

30 The invention also provides the use of polynucleotide shuffling to shuffle a population of protein variants, such as taxonomically-related, structurally-related, and/or functionally-related enzymes and/or mutated variants thereof to create and identify advantageous novel polypeptides, such as enzymes having
35 altered properties of catalysis, temperature profile, stability, oxidation resistance, or other desired feature which can be selected for. Methods suitable for molecular evolution and

directed molecular evolution are provided. Methods to focus selection pressure(s) upon specific portions of polynucleotides (such as a segment of a coding region) are provided.

The invention also provides a method suitable for 5 shuffling polynucleotide sequences for generating gene therapy vectors and replication-defective gene therapy constructs, such as may be used for human gene therapy, including but not limited to vaccination vectors for DNA-based vaccination, as well as anti-neoplastic gene therapy and other gene therapy formats.

10 The invention provides a method for generating an enhanced green fluorescent protein (GFP) and polynucleotides encoding same, comprising performing DNA shuffling on a GFP encoding expression vector and selecting or screening for variants having an enhanced desired property, such as enhanced 15 fluorescence. In a variation, an embodiment comprises a step of error-prone or mutagenic amplification, propagation in a mutator strain (e.g., a host cell having a hypermutational phenotype; mut^L, etc.; yeast strains such as those described in Klein (1995) 20 Progr. Nucl. Acid Res. Mol. Biol. 51: 217, incorporated herein by reference), chemical mutagenesis, or site-directed mutagenesis. In an embodiment, the enhanced GFP protein comprises a point mutation outside the chromophore region (amino acids 64-69), preferably in the region from amino acid 100 to 25 amino acid 173, with specific preferred embodiments at residue 100, 154, and 164; typically, the mutation is a substitution mutation, such as F100S, M154T or V164A. In an embodiment, the mutation substitutes a hydrophilic residue for a hydrophobic residue. In an embodiment, multiple mutations are present in the 30 enhanced GFP protein and its encoding polynucleotide. The invention also provides the use of such an enhanced GFP protein, such as for a diagnostic reporter for assays and high throughput screening assays and the like.

The invention also provides for improved embodiments for performing in vitro sequence shuffling. In one aspect, the 35 improved shuffling method includes the addition of at least one additive which enhances the rate or extent of reannealing or recombination of related-sequence polynucleotides. In an

embodiment, the additive is polyethylene glycol (PEG), typically added to a shuffling reaction to a final concentration of 0.1 to 25 percent, often to a final concentration of 2.5 to 15 percent, to a final concentration of about 10 percent. In an embodiment,
5 the additive is dextran sulfate, typically added to a shuffling reaction to a final concentration of 0.1 to 25 percent, often at about 10 percent. In an embodiment, the additive is an agent which reduces sequence specificity of reannealing and promotes promiscuous hybridization and/or recombination in vitro. In an
10 alternative embodiment, the additive is an agent which increases sequence specificity of reannealing and promotes high fidelity hybridization and/or recombination in vitro. Other long-chain polymers which do not interfere with the reaction may also be used (e.g., polyvinylpyrrolidone, etc.).

15 In one aspect, the improved shuffling method includes the addition of at least one additive which is a cationic detergent. Examples of suitable cationic detergents include but are not limited to: cetyltrimethylammonium bromide (CTAB), dodecyltrimethylammonium bromide (DTAB), and tetramethylammonium
20 chloride (TMAC), and the like.

In one aspect, the improved shuffling method includes the addition of at least one additive which is a recombinogenic protein that catalyzes or non-catalytically enhances homologous pairing and/or strand exchange in vitro. Examples of suitable recombinogenic proteins include but are not limited to: *E. coli* recA protein, the T4 uvsX protein, the recI protein from *Ustilago maydis*, other recA family recombinases from other species, single strand binding protein (SSB), ribonucleoprotein A1, and the like. Shuffling can be used to improve one or more properties of a recombinogenic protein; for example, mutant sequences encoding recA can be shuffled and improved heat-stable variants selected by recursive sequence recombination.

Non-specific (general recombination) recombinases such as Topoisomerase I, Topoisomerase II (Tse et al. (1980) J. Biol. Chem. 255: 5560; Trask et al. (1984) EMBO J. 3: 671, incorporated herein by reference) and the like can be used to catalyze in vitro recombination reactions to shuffle a plurality of related

sequence polynucleotide species by the recursive methods of the invention.

In one aspect, the improved shuffling method includes the addition of at least one additive which is an enzyme having 5 an exonuclease activity which is active at removing non-templated nucleotides introduced at 3' ends of product polynucleotides in shuffling amplification reactions catalyzed by a non-proofreading polymerase. An example of a suitable enzyme having an exonuclease activity includes but is not limited to Pfu 10 polymerase. Other suitable polymerases include, but are not limited to: *Thermus flavus* DNA polymerase (Tfl)
Thermus thermophilus DNA polymerase (Tth)
Thermococcus litoralis DNA polymerase (Tli, Vent)
Pyrococcus Woeselii DNA polymerase (Pwo)
15 *Thermotoga maritima* DNA polymerase (UltMa)
Thermus brockianus DNA polymerase (Thermozyme)
Pyrococcus furiosus DNA polymerase (Pfu)
Thermococcus sp. DNA polymerase (9'Nm)
Pyrococcus sp. DNA polymerase ('Deep Vent')
20 Bacteriophage T4 DNA polymerase
Bacteriophage T7 DNA polymerase
E. coli DNA polymerase I (native and Klenow)
E. coli DNA polymerase III.

In an aspect, the improved shuffling method comprises 25 the modification wherein at least one cycle of amplification (i.e., extension with a polymerase) of reannealed fragmented library member polynucleotides is conducted under conditions which produce a substantial fraction, typically at least 20 percent or more, of incompletely extended amplification products.
30 The amplification products, including the incompletely extended amplification products are denatured and subjected to at least one additional cycle of reannealing and amplification. This variation, wherein at least one cycle of reannealing and amplification provides a substantial fraction of incompletely 35 extended products, is termed "stuttering" and in the subsequent amplification round the incompletely extended products reanneal

to and prime extension on different sequence-related template species.

In an aspect, the improved shuffling method comprises the modification wherein at least one cycle of amplification is 5 conducted using a collection of overlapping single-stranded DNA fragments of varying lengths corresponding to a first polynucleotide species or set of related-sequence polynucleotide species, wherein each overlapping fragment can each hybridize to and prime polynucleotide chain extension from a second 10 polynucleotide species serving as a template, thus forming sequence-recombined polynucleotides, wherein said sequence-recombined polynucleotides comprise a portion of at least one first polynucleotide species with an adjacent portion of the second polynucleotide species which serves as a template. In a 15 variation, the second polynucleotide species serving as a template contains uracil (i.e., a Kunkel-type template) and is substantially non-replicable in cells. This aspect of the invention can also comprise at least two recursive cycles of this variation.

20 In an embodiment, PCR can be conducted wherein the nucleotide mix comprises a nucleotide species having uracil as the base. The PCR product(s) can then be fragmented by digestion with UDG-glycosylase which produces strand breaks. The fragment size can be controlled by the fraction of uracil-containing NTP 25 in the PCR mix.

In an aspect, the improved shuffling method comprises the modification wherein at least one cycle of amplification is conducted with an additive or polymerase in suitable conditions which promote template switching. In an embodiment where Taq 30 polymerase is employed for amplification, addition of recA or other polymerases (e.g., viral polymerases, reverse transcriptase) enhances template switching. Template-switching can also be increased by increasing the DNA template concentration, among other means known by those skilled in the 35 art.

In an embodiment of the general method, libraries of sequence-recombined polynucleotides are generated from sequence-

related polynucleotides which are naturally-occurring genes or alleles of a gene. In this aspect, at least two naturally-occurring genes and/or alleles which comprise regions of at least 50 consecutive nucleotides which have at least 70 percent sequence identity, preferably at least 90 percent sequence identity, are selected from a pool of gene sequences, such as by hybrid selection or via computerized sequence analysis using sequence data from a database. In an aspect, at least three naturally-occurring genes and/or alleles which comprise regions of at least 50 consecutive nucleotides which have at least 70 percent sequence identity, preferably at least 90 percent sequence identity, are selected from a pool of gene sequences, such as by hybrid selection or via computerized sequence analysis using sequence data from a database. The selected sequences are obtained as polynucleotides, either by cloning or via DNA synthesis, and shuffled by any of the various embodiments of the invention.

In an embodiment of the invention, multi-pool shuffling is performed. Shuffling of multiple pools of polynucleotide sequences allows each separate pool to generate a different combinatorial solution to produce the desired property. In this variation, the pool of parental polynucleotides sequences (or any subsequent shuffled library or selected pool of library members) is subdivided (or segregated) into two or more discrete pools of sequences and are separately subjected to one or more rounds of recursive sequence recombination and selection (or screening). If desired, optionally, selected library members from each separate pool may be recombined (integrated) in latter rounds of shuffling. Alternatively, multiple separate parental pools may be used. Inbreeding, wherein selected (or screened) library members within a pool are crossed with each other by the recursive sequence recombination methods of the invention, can be performed, alone or in combination with outbreeding, wherein library members of different pools are crossed with each other by the recursive sequence recombination methods of the invention.

In an embodiment of the invention, the method comprises the further step of removing non-shuffled products (e.g.,

parental sequences) from sequence-recombined polynucleotides produced by any of the disclosed shuffling methods. Non-shuffled products can be removed or avoided by performing amplification with: (1) a first PCR primer which hybridizes to a first 5 parental polynucleotide species but does not substantially hybridize to a second parental polynucleotide species, and (2) a second PCR primer which hybridizes to a second parental polynucleotide species but does not substantially hybridize to the first parental polynucleotide species, such that 10 amplification occurs from templates comprising the portion of the first parental sequence which hybridizes to the first PCR primer and also comprising the portion of the second parental sequence which hybridizes to the second PCR primer, thus only sequence-recombined polynucleotides are amplified.

15 The invention also provides for alternative embodiments for performing in vivo sequence shuffling. In one aspect, the alternative shuffling method includes the use of inter-plasmidic recombination, wherein libraries of sequence-recombined polynucleotide sequences are obtained by genetic recombination 20 in vivo of compatible or non-compatible multicopy plasmids inside suitable host cells. When non-compatible plasmids are used, typically each plasmid type has a distinct selectable marker and selection for retention of each desired plasmid type is applied. The related-sequence polynucleotide sequences to be recombined 25 are separately incorporated into separately replicable multicopy vectors, typically bacterial plasmids each having a distinct and separately selectable marker gene (e.g., a drug resistance gene). Suitable host cells are transformed with both species of plasmid and cells expressing both selectable marker genes are selected 30 and sequence-recombined sequences are recovered and can be subjected to additional rounds of shuffling by any of the means described herein.

In one aspect, the alternative shuffling method includes the use of intra-plasmidic recombination, wherein 35 libraries of sequence-recombined polynucleotide sequences are obtained by genetic recombination in vivo of direct or inverted sequence repeats located on the same plasmid. In a variation,

the sequences to be recombined are flanked by site-specific recombination sequences and the polynucleotides are present in a site-specific recombination system, such as an integron (Hall and Collins (1995) Mol. Microbiol. **15**: 593, incorporated herein by reference) and can include insertion sequences, transposons (e.g., IS1), and the like. Introns have a low rate of natural mobility and can be used as mobile genetic elements both in prokaryotes and eukaryotes. Shuffling can be used to improve the performance of mobile genetic elements. These high frequency recombination vehicles can be used for the rapid optimization of large sequences via transfer of large sequence blocks. Recombination between repeated, interspersed, and diverged DNA sequences, also called "homeologous" sequences, is typically suppressed in normal cells. However, in MutL and MutS cells, this suppression is relieved and the rate of intrachromosomal recombination is increased (Petit et al. (1996) Genetics **129**: 327, incorporated herein by reference).

In an aspect of the invention, mutator strains of host cells are used to enhance recombination of more highly mismatched sequence-related polynucleotides. Bacterial strains such as MutL, MutS, MutT, or MutH or other cells expressing the Mut proteins (XL-1red; Stratagene, San Diego, CA) can be used as host cells for shuffling of sequence-related polynucleotides by in vivo recombination. Other mutation-prone host cell types can also be used, such as those having a proofreading-defective polymerase (Foster et al. (1995) Proc. Natl. Acad. Sci. (U.S.A.) **92**: 7951, incorporated herein by reference). Mutator strains of yeast can be used, as can hypermutational mammalian cells, including ataxia telangiectasia cells, such as described in Luo et al. (1996) J. Biol. Chem. **271**: 4497, incorporated herein by reference.

Other in vivo and in vitro mutagenic formats can be employed, including administering chemical or radiological mutagens to host cells. Examples of such mutagens include but are not limited to: MNU, ENU, MNNG, nitrosourea, BuDR, and the like. Ultraviolet light can also be used to generate mutations and/or to enhance the rate of recombination, such as by

irradiation of cells used to enhance *in vivo* recombination. Ionizing radiation and clastogenic agents can also be used to enhance mutational frequency and/or to enhance recombination and/or to effect polynucleotide fragmentation.

5

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 is a schematic diagram comparing mutagenic shuffling over error-prone PCR; (a) the initial library; (b) pool of selected sequences in first round of affinity selection; (d) 10 *in vitro* recombination of the selected sequences ('shuffling'); (f) pool of selected sequences in second round of affinity selection after shuffling; (c) error-prone PCR; (e) pool of selected sequences in second round of affinity selection after error-prone PCR.

15 Figure 2 illustrates the reassembly of a 1.0 kb LacZ alpha gene fragment from 10-50 bp random fragments. (a) Photograph of a gel of PCR amplified DNA fragment having the LacZ alpha gene. (b) Photograph of a gel of DNA fragments after digestion with DNaseI. (c) Photograph of a gel of DNA fragments 20 of 10-50 bp purified from the digested LacZ alpha gene DNA fragment; (d) Photograph of a gel of the 10-50 bp DNA fragments after the indicated number of cycles of DNA reassembly; (e) Photograph of a gel of the recombination mixture after amplification by PCR with primers.

25 Figure 3 is a schematic illustration of the LacZ alpha gene stop codon mutants and their DNA sequences. The boxed regions are heterologous areas, serving as markers. The stop codons are located in smaller boxes or underlined. "+" indicates a wild-type gene and "-" indicates a mutated area in the gene.

30 Figure 4 is a schematic illustration of the introduction or spiking of a synthetic oligonucleotide into the reassembly process of the LacZ alpha gene.

35 Figure 5 illustrates the regions of homology between a murine IL1-B gene (M) and a human IL1-B gene (H) with *E. coli* codon usage. Regions of heterology are boxed. The "_|_" indicate crossovers obtained upon the shuffling of the two genes.

Figure 6 is a schematic diagram of the antibody CDR shuffling model system using the scFv of anti-rabbit IgG antibody (A10B).

Figure 7 (panels A-D) illustrates the observed frequency of occurrence of certain combinations of CDRs in the shuffled DNA of the scFv. Panel (A) shows the length and mutagenesis rate of all six synthetic CDRs. Panel (B) shows library construction by shuffling scFv with all six CDRs. Panel (C) shows CDR insertion determined by PCR with primers for native CDRs. Panel (D) shows insertion rates and distributions of synthetic CDR insertions.

Figure 8 illustrates the improved avidity of the scFv anti-rabbit antibody after DNA shuffling and each cycle of selection.

Figure 9 schematically portrays pBR322-Sfi-BL-LA-Sfi and in vivo intraplasmidic recombination via direct repeats, as well as the rate of generation of ampicillin-resistant colonies by intraplasmidic recombination reconstituting a functional beta-lactamase gene.

Figure 10 schematically portrays pBR322-Sfi-2Bla-Sfi and in vivo intraplasmidic recombination via direct repeats, as well as the rate of generation of ampicillin-resistant colonies by intraplasmidic recombination reconstituting a functional beta-lactamase gene.

Figure 11 illustrates the method for testing the efficiency of multiple rounds of homologous recombination after the introduction of polynucleotide fragments into cells for the generation of recombinant proteins.

Figure 12 schematically portrays generation of a library of vectors by shuffling cassettes at the following loci: promoter, leader peptide, terminator, selectable drug resistance gene, and origin of replication. The multiple parallel lines at each locus represent the multiplicity of cassettes for that cassette.

Figure 13 schematically shows some examples of cassettes suitable at various loci for constructing prokaryotic vector libraries by shuffling.

Figure 14 shows the prokaryotic GFP expression vector PBAD-GFP (5,371 bp) was derived from pBAD18 (Guzman et al. (1995) J. Bacteriol. 177: 4121). The eukaryotic GFP expression vector Alpha+GFP (7,591 bp) was derived from the vector Alpha+ (Whitehorn et al. (1995) Bio/Technology 13: 1215).

Figure 15A and 15B show comparison of the fluorescence of different GFP constructs in whole *E. coli* cells. Compared are the 'Clontech' construct which contains a 24 amino acid N-terminal extension, the Affymax wildtype construct ('wt', with improved codon usage), and the mutants obtained after 2 and after 3 cycles of sexual PCR and selection ('cycle 2', 'cycle 3'). The 'Clontech' construct was induced with IPTG, whereas the other constructs were induced with arabinose. All samples were assayed at equal OD₆₀₀. Figure 15A shows fluorescence spectra indicating that the whole cell fluorescence signal from the 'wt' construct is 2.8-fold greater than from the 'Clontech' construct. The signal of the 'cycle 3' mutant is 16-fold increased over the Affymax 'wt', and 45-fold over the 'Clontech' wt construct. Figure 15B is a comparison of excitation spectra of GFP constructs in *E. coli*. The peak excitation wavelengths are unaltered by the mutations that were selected.

Figure 16 shows SDS-PAGE analysis of relative GFP protein expression levels. Panel (a): 12% Tris-Glycine SDS-PAGE analysis (Novex, Encinitas, CA) of equal amounts (OD₆₀₀) Of whole *E. coli* cells expressing the wildtype, the cycle 2 mutant or the cycle 3 mutant of GFP. Stained with Coomassie Blue. GFP (27 kD) represents about 75% of total protein, and the selection did not increase the expression level. Panel (b) 12% Tris-Glycine SDS-PAGE analysis (Novex, Encinitas, CA) of equal amounts (OD₆₀₀) of *E. coli* fractions. Lane 1: Pellet of lysed cells expressing wt GFP; lane 2: Supernatant of lysed cells expressing wt GFP. Most of the wt GFP is in inclusion bodies; lane 3: Pellet of lysed cells expressing cycle 3 mutant GFP; lane 4: Supernatant of lysed cells expressing cycle 3 mutant GFP. Most of the wt GFP is soluble. The GFP that ends up in inclusion bodies does not contain the chromophore, since there is no detectable fluorescence in this fraction.

Figure 17 shows mutation analysis of the cycle 2 and cycle 3 mutants versus wildtype GFP. Panel (A) shows that the mutations are spread out rather than clustered near the tripeptide chromophore. Mutations F100S, M154T, and V164A involve the replacement of hydrophobic residues with more hydrophilic residues. The increased hydrophilicity may help guide the protein into a native folding pathway rather than toward aggregation and inclusion body formation. Panel (B) shows a restriction map indicating the chromophore region and positions of introduced mutations.

Figure 18A and 18B show comparison of CHO cells expressing different GFP proteins. Figure 18A is a FACS analysis of clones of CHO cells expressing different GFP mutants. Figure 18B shows fluorescence spectroscopy of clones of CHO cells expressing different GFP mutants.

Figure 19 shows enhancement of resistance to arsenate toxicity as a result of shuffling the pGJ103 plasmid containing the arsenate detoxification pathway operon.

Figure 20 schematically shows the generation of combinatorial libraires using synthetic or naturally-occurring intron sequences as the basis for recombining a plurality of exons species which can lack sequence identity (as exemplified by random sequence exons), wherein homologous and/or site-specific recombination occurs between intron sequences of distinct library members.

Figure 21 schematically shows variations of the method for shuffling exons. The numbers refer to reading frames, as demonstrated in panel (A). Panel (B) shows the various classes of intron and exon relative to their individual splice frames. Panel (C) provides an example of a naturally-occurring gene (immunoglobulin V genes) suitable for shuffling. Panels D through F shows how multiple exons can be concatemerized via PCR using primers which span intron segments, so that proper splicing frames are retained, if desired. Panel (G) exemplifies the exon shuffling process (IG: immunoglobulin exon; IFN: interferon exon)

Figure 22 schematically shows an exon splicing frame diagram for several human genes, showing that preferred units for

shuffling exons begin and end in the same splicing frame, such that a splicing module (or shuffling exon) can comprise multiple naturally-occurring exons but typically has the same splicing frame at each end.

5 Figure 23 schematically shows how partial PCR extension (stuttering) can be used to provide recursive sequence recombination (shuffling) resulting in a library of chimeras representing multiple crossovers.

10 Figure 24 shows how stuttering can be used to shuffle a wild-type sequence with a multiply mutated sequence to generate an optimal set of mutations via shuffling.

15 Figure 25 schematically shows plasmid-plasmid recombination by electroporation of a cell population representing multiple plasmid species, present initially as a single plasmid species per cell prior to electroporation and multiple plasmid species per cell suitable for in vivo recombination subsequent to electroporation of the cell population.

20 Figure 26 shows plasmid-plasmid recombination.

Figure 27 shows plasmid-virus recombination.

Figure 28 shows virus-virus recombination.

Figure 29 shows plasmid-chromosome recombination.

Figure 30 shows conjugation-mediated recombination.

25 Figure 31 shows Ab-phage recovery rate versus selection cycle. Shuffling was applied after selection rounds two to eight. Total increase is 440-fold.

30 Figure 32 shows binding specificity after ten selection rounds, including two rounds of backcrossing. ELISA signal of different Ab-phage clones for eight human protein targets.

Figure 33 shows Ab-phage recovery versus mutagenesis method.

DESCRIPTION OF THE PREFERRED EMBODIMENTS

35 The present invention relates to a method for nucleic acid molecule reassembly after random fragmentation and its application to mutagenesis of DNA sequences. Also described is a method for the production of nucleic acid fragments encoding

mutant proteins having enhanced biological activity. In particular, the present invention also relates to a method of repeated cycles of mutagenesis, nucleic acid shuffling and selection which allow for the creation of mutant proteins having 5 enhanced biological activity.

The present invention is directed to a method for generating a very large library of DNA, RNA or protein mutants; in embodiments where a metabolic enzyme or multicomponent pathway is subjected to shuffling, a library can compose the resultant 10 metabolites in addition to a library of the shuffled enzyme(s). This method has particular advantages in the generation of related DNA fragments from which the desired nucleic acid fragment(s) may be selected. In particular the present invention 15 also relates to a method of repeated cycles of mutagenesis, homologous recombination and selection which allow for the creation of mutant proteins having enhanced biological activity.

However, prior to discussing this invention in further detail, the following terms will first be defined.

20 Definitions

As used herein, the following terms have the following meanings:

The term "DNA reassembly" is used when recombination occurs between identical sequences.

25 By contrast, the term "DNA shuffling" is used herein to indicate recombination between substantially homologous but non-identical sequences, in some embodiments DNA shuffling may involve crossover via nonhomologous recombination, such as via cre/lox and/or flp/frt systems and the like, such that 30 recombination need not require substantially homologous polynucleotide sequences. Homologous and non-homologous recombination formats can be used, and, in some embodiments, can generate molecular chimeras and/or molecular hybrids of substantially dissimilar sequences.

35 The term "amplification" means that the number of copies of a nucleic acid fragment is increased.

The term "identical" or "identity" means that two nucleic acid sequences have the same sequence or a complementary sequence. Thus, "areas of identity" means that regions or areas of a nucleic acid fragment or polynucleotide are identical or 5 complementary to another polynucleotide or nucleic acid fragment.

The term "corresponds to" is used herein to mean that a polynucleotide sequence is homologous (i.e., is identical, not strictly evolutionarily related) to all or a portion of a reference polynucleotide sequence, or that a polypeptide sequence 10 is identical to a reference polypeptide sequence. In contradistinction, the term "complementary to" is used herein to mean that the complementary sequence is homologous to all or a portion of a reference polynucleotide sequence. For illustration, the nucleotide sequence "TATAC" corresponds to a reference 15 sequence "TATAC" and is complementary to a reference sequence "GTATA".

The following terms are used to describe the sequence relationships between two or more polynucleotides: "reference sequence", "comparison window", "sequence identity", "percentage 20 of sequence identity", and "substantial identity". A "reference sequence" is a defined sequence used as a basis for a sequence comparison; a reference sequence may be a subset of a larger sequence, for example, as a segment of a full-length cDNA or gene sequence given in a sequence listing, such as a polynucleotide 25 sequence of Fig. 1 or Fig. 2(b), or may comprise a complete cDNA or gene sequence. Generally, a reference sequence is at least 20 nucleotides in length, frequently at least 25 nucleotides in length, and often at least 50 nucleotides in length. Since two polynucleotides may each (1) comprise a sequence (i.e., a portion 30 of the complete polynucleotide sequence) that is similar between the two polynucleotides, and (2) may further comprise a sequence that is divergent between the two polynucleotides, sequence comparisons between two (or more) polynucleotides are typically performed by comparing sequences of the two polynucleotides over 35 a "comparison window" to identify and compare local regions of sequence similarity.

A "comparison window", as used herein, refers to a conceptual segment of at least 20 contiguous nucleotide positions wherein a polynucleotide sequence may be compared to a reference sequence of at least 20 contiguous nucleotides and wherein the portion of the polynucleotide sequence in the comparison window may comprise additions or deletions (i.e., gaps) of 20 percent or less as compared to the reference sequence (which does not comprise additions or deletions) for optimal alignment of the two sequences. Optimal alignment of sequences for aligning a comparison window may be conducted by the local homology algorithm of Smith and Waterman (1981) Adv. Appl. Math. 2: 482, by the homology alignment algorithm of Needleman and Wunsch (1970) J. Mol. Biol. 48: 443, by the search for similarity method of Pearson and Lipman (1988) Proc. Natl. Acad. Sci. (U.S.A.) 85: 2444, by computerized implementations of these algorithms (GAP, BESTFIT, FASTA, and TFASTA in the Wisconsin Genetics Software Package Release 7.0, Genetics Computer Group, 575 Science Dr., Madison, WI), or by inspection, and the best alignment (i.e., resulting in the highest percentage of homology over the comparison window) generated by the various methods is selected.

The term "sequence identity" means that two polynucleotide sequences are identical (i.e., on a nucleotide-by-nucleotide basis) over the window of comparison. The term "percentage of sequence identity" is calculated by comparing two optimally aligned sequences over the window of comparison, determining the number of positions at which the identical nucleic acid base (e.g., A, T, C, G, U, or I) occurs in both sequences to yield the number of matched positions, dividing the number of matched positions by the total number of positions in the window of comparison (i.e., the window size), and multiplying the result by 100 to yield the percentage of sequence identity. The terms "substantial identity" as used herein denotes a characteristic of a polynucleotide sequence, wherein the polynucleotide comprises a sequence that has at least 80 percent sequence identity, preferably at least 85 percent identity and often 90 to 95 percent sequence identity, more usually at least 99 percent sequence identity as compared to a reference sequence.

over a comparison window of at least 20 nucleotide positions, frequently over a window of at least 25-50 nucleotides, wherein the percentage of sequence identity is calculated by comparing the reference sequence to the polynucleotide sequence which may 5 include deletions or additions which total 20 percent or less of the reference sequence over the window of comparison.

Conservative amino acid substitutions refer to the interchangeability of residues having similar side chains. For example, a group of amino acids having aliphatic side chains is 10 glycine, alanine, valine, leucine, and isoleucine; a group of amino acids having aliphatic-hydroxyl side chains is serine and threonine; a group of amino acids having amide-containing side chains is asparagine and glutamine; a group of amino acids having aromatic side chains is phenylalanine, tyrosine, and tryptophan; 15 a group of amino acids having basic side chains is lysine, arginine, and histidine; and a group of amino acids having sulfur-containing side chains is cysteine and methionine. Preferred conservative amino acids substitution groups are: valine-leucine-isoleucine, phenylalanine-tyrosine, lysine-20 arginine, alanine-valine, and asparagine-glutamine.

The term "homologous" or "homeologous" means that one single-stranded nucleic acid sequence may hybridize to a complementary single-stranded nucleic acid sequence. The degree of hybridization may depend on a number of factors including the 25 amount of identity between the sequences and the hybridization conditions such as temperature and salt concentration as discussed later. Preferably the region of identity is greater than about 5 bp, more preferably the region of identity is greater than 10 bp.

30 The term "heterologous" means that one single-stranded nucleic acid sequence is unable to hybridize to another single-stranded nucleic acid sequence or its complement. Thus areas of heterology means that nucleic acid fragments or polynucleotides have areas or regions in the sequence which are unable to 35 hybridize to another nucleic acid or polynucleotide. Such regions or areas are, for example, areas of mutations.

The term "cognate" as used herein refers to a gene sequence that is evolutionarily and functionally related between species. For example but not limitation, in the human genome, the human CD4 gene is the cognate gene to the mouse CD4 gene, 5 since the sequences and structures of these two genes indicate that they are highly homologous and both genes encode a protein which functions in signaling T cell activation through MHC class II-restricted antigen recognition.

The term "wild-type" means that the nucleic acid fragment 10 does not comprise any mutations. A "wild-type" protein means that the protein will be active at a level of activity found in nature and typically will comprise the amino acid sequence found in nature. In an aspect, the term "wild type" or "parental sequence" can indicate a starting or reference sequence prior to 15 a manipulation of the invention.

The term "related polynucleotides" means that regions or areas of the polynucleotides are identical and regions or areas of the polynucleotides are heterologous.

The term "chimeric polynucleotide" means that the 20 polynucleotide comprises regions which are wild-type and regions which are mutated. It may also mean that the polynucleotide comprises wild-type regions from one polynucleotide and wild-type regions from another related polynucleotide.

The term "cleaving" means digesting the polynucleotide with 25 enzymes or breaking the polynucleotide, or generating partial length copies of a parent sequence(s) via partial PCR extension, PCR stuttering, differential fragment amplification, or other means of producing partial length copies of one or more parental sequences.

The term "population" as used herein means a collection of 30 components such as polynucleotides, nucleic acid fragments or proteins. A "mixed population" means a collection of components which belong to the same family of nucleic acids or proteins (i.e. are related) but which differ in their sequence (i.e. are 35 not identical) and hence in their biological activity.

The term "specific nucleic acid fragment" means a nucleic acid fragment having certain end points and having a certain

nucleic acid sequence. Two nucleic acid fragments wherein one nucleic acid fragment has the identical sequence as a portion of the second nucleic acid fragment but different ends comprise two different specific nucleic acid fragments.

5 The term "mutations" means changes in the sequence of a wild-type nucleic acid sequence or changes in the sequence of a peptide. Such mutations may be point mutations such as transitions or transversions. The mutations may be deletions, insertions or duplications.

10 In the polypeptide notation used herein, the lefthand direction is the amino terminal direction and the righthand direction is the carboxy-terminal direction, in accordance with standard usage and convention. Similarly, unless specified otherwise, the lefthand end of single-stranded polynucleotide sequences is the 5' end; the lefthand direction of double-stranded polynucleotide sequences is referred to as the 5' direction. The direction of 5' to 3' addition of nascent RNA transcripts is referred to as the transcription direction; sequence regions on the DNA strand having the same sequence as 15 the RNA and which are 5' to the 5' end of the RNA transcript are referred to as "upstream sequences"; sequence regions on the DNA strand having the same sequence as the RNA and which are 3' to the 3' end of the coding RNA transcript are referred to as 20 "downstream sequences".

25 The term "naturally-occurring" as used herein as applied to an object refers to the fact that an object can be found in nature. For example, a polypeptide or polynucleotide sequence that is present in an organism (including viruses) that can be isolated from a source in nature and which has not been 30 intentionally modified by man in the laboratory is naturally-occurring. Generally, the term naturally-occurring refers to an object as present in a non-pathological (undiseased) individual, such as would be typical for the species.

The term "agent" is used herein to denote a chemical 35 compound, a mixture of chemical compounds, an array of spatially localized compounds (e.g., a VLSIPS peptide array, polynucleotide array, and/or combinatorial small molecule array), a biological

macromolecule, a bacteriophage peptide display library, a bacteriophage antibody (e.g., scFv) display library, a polysome peptide display library, or an extract made from biological materials such as bacteria, plants, fungi, or animal
5 (particularly mammalian) cells or tissues. Agents are evaluated for potential activity as antineoplastics, anti-inflammatories, or apoptosis modulators by inclusion in screening assays described hereinbelow. Agents are evaluated for potential activity as specific protein interaction inhibitors (i.e., an
10 agent which selectively inhibits a binding interaction between two predetermined polypeptides but which does not substantially interfere with cell viability) by inclusion in screening assays described hereinbelow.

As used herein, "substantially pure" means an object species is the predominant species present (i.e., on a molar basis it is more abundant than any other individual macromolecular species in the composition), and preferably a substantially purified fraction is a composition wherein the object species comprises at least about 50 percent (on a molar
15 basis) of all macromolecular species present. Generally, a substantially pure composition will comprise more than about 80 to 90 percent of all macromolecular species present in the composition. Most preferably, the object species is purified to essential homogeneity (contaminant species cannot be detected in
20 the composition by conventional detection methods) wherein the composition consists essentially of a single macromolecular species. Solvent species, small molecules (<500 Daltons), and elemental ion species are not considered macromolecular species.
25

As used herein the term "physiological conditions"
30 refers to temperature, pH, ionic strength, viscosity, and like biochemical parameters which are compatible with a viable organism, and/or which typically exist intracellularly in a viable cultured yeast cell or mammalian cell. For example, the intracellular conditions in a yeast cell grown under typical
35 laboratory culture conditions are physiological conditions. Suitable in vitro reaction conditions for in vitro transcription cocktails are generally physiological conditions. In general,

in *vitro* physiological conditions comprise 50-200 mM NaCl or KCl, pH 6.5-8.5, 20-45°C and 0.001-10 mM divalent cation (e.g., Mg⁺⁺, Ca⁺⁺); preferably about 150 mM NaCl or KCl, pH 7.2-7.6, 5 mM divalent cation, and often include 0.01-1.0 percent nonspecific protein (e.g., BSA). A non-ionic detergent (Tween, NP-40, Triton X-100) can often be present, usually at about 0.001 to 2%, typically 0.05-0.2% (v/v). Particular aqueous conditions may be selected by the practitioner according to conventional methods. For general guidance, the following buffered aqueous conditions may be applicable: 10-250 mM NaCl, 5-50 mM Tris HCl, pH 5-8, with optional addition of divalent cation(s) and/or metal chelators and/or nonionic detergents and/or membrane fractions and/or antifoam agents and/or scintillants.

Specific hybridization is defined herein as the formation of hybrids between a first polynucleotide and a second polynucleotide (e.g., a polynucleotide having a distinct but substantially identical sequence to the first polynucleotide), wherein the first polynucleotide preferentially hybridizes to the second polynucleotide under stringent hybridization conditions wherein substantially unrelated polynucleotide sequences do not form hybrids in the mixture.

As used herein, the term "single-chain antibody" refers to a polypeptide comprising a V_H domain and a V_L domain in polypeptide linkage, generally linked via a spacer peptide (e.g., [Gly-Gly-Gly-Ser]_x), and which may comprise additional amino acid sequences at the amino- and/or carboxy- termini. For example, a single-chain antibody may comprise a tether segment for linking to the encoding polynucleotide. As an example, a scFv is a single-chain antibody. Single-chain antibodies are generally proteins consisting of one or more polypeptide segments of at least 10 contiguous amino acids substantially encoded by genes of the immunoglobulin superfamily (e.g., see The Immunoglobulin Gene Superfamily, A.F. Williams and A.N. Barclay, in Immunoglobulin Genes, T. Honjo, F.W. Alt, and T.H. Rabbitts, eds., (1989) Academic Press: San Diego, CA, pp.361-387, which is incorporated herein by reference), most frequently encoded by a rodent, non-human primate, avian, porcine, bovine, ovine, goat,

or human heavy chain or light chain gene sequence. A functional single-chain antibody generally contains a sufficient portion of an immunoglobulin superfamily gene product so as to retain the property of binding to a specific target molecule, typically a 5 receptor or antigen (epitope).

As used herein, the term "complementarity-determining region" and "CDR" refer to the art-recognized term as exemplified by the Kabat and Chothia CDR definitions also generally known as hypervariable regions or hypervariable loops (Chothia and Lesk 10 (1987) J. Mol. Biol. **196**: 901; Chothia et al. (1989) Nature **342**: 877; E.A. Kabat et al., Sequences of Proteins of Immunological Interest (National Institutes of Health, Bethesda, MD) (1987); and Tramontano et al. (1990) J. Mol. Biol. **215**: 175). Variable 15 region domains typically comprise the amino-terminal approximately 105-115 amino acids of a naturally-occurring immunoglobulin chain (e.g., amino acids 1-110), although variable domains somewhat shorter or longer are also suitable for forming single-chain antibodies.

An immunoglobulin light or heavy chain variable region 20 consists of a "framework" region interrupted by three hypervariable regions, also called CDR's. The extent of the framework region and CDR's have been precisely defined (see, "Sequences of Proteins of Immunological Interest," E. Kabat et al., 4th Ed., U.S. Department of Health and Human Services, 25 Bethesda, MD (1987)). The sequences of the framework regions of different light or heavy chains are relatively conserved within a species. As used herein, a "human framework region" is a framework region that is substantially identical (about 85% or more, usually 90-95% or more) to the framework region of a 30 naturally occurring human immunoglobulin. The framework region of an antibody, that is the combined framework regions of the constituent light and heavy chains, serves to position and align the CDR's. The CDR's are primarily responsible for binding to an epitope of an antigen.

35 As used herein, the term "variable segment" refers to a portion of a nascent peptide which comprises a random, pseudorandom, or defined kernal sequence. A variable segment can

comprise both variant and invariant residue positions, and the degree of residue variation at a variant residue position may be limited; both options are selected at the discretion of the practitioner. Typically, variable segments are about 5 to 20 5 amino acid residues in length (e.g., 8 to 10), although variable segments may be longer and may comprise antibody portions or receptor proteins, such as an antibody fragment, a nucleic acid binding protein, a receptor protein, and the like.

As used herein, "random peptide sequence" refers to an 10 amino acid sequence composed of two or more amino acid monomers and constructed by a stochastic or random process. A random peptide can include framework or scaffolding motifs, which may comprise invariant sequences.

As used herein "random peptide library" refers to a set 15 of polynucleotide sequences that encodes a set of random peptides, and to the set of random peptides encoded by those polynucleotide sequences, as well as the fusion proteins containing those random peptides.

As used herein, the term "pseudorandom" refers to a set 20 of sequences that have limited variability, so that for example the degree of residue variability at one position is different than the degree of residue variability at another position, but any pseudorandom position is allowed some degree of residue variation, however circumscribed.

As used herein, the term "defined sequence framework" 25 refers to a set of defined sequences that are selected on a nonrandom basis, generally on the basis of experimental data or structural data; for example, a defined sequence framework may comprise a set of amino acid sequences that are predicted to form 30 a β -sheet structure or may comprise a leucine zipper heptad repeat motif, a zinc-finger domain, among other variations. A "defined sequence kernal" is a set of sequences which encompass 35 a limited scope of variability. Whereas (1) a completely random 10-mer sequence of the 20 conventional amino acids can be any of (20)¹⁰ sequences, and (2) a pseudorandom 10-mer sequence of the 20 conventional amino acids can be any of (20)¹⁰ sequences but will exhibit a bias for certain residues at certain positions

and/or overall, (3) a defined sequence kernal is a subset of sequences which is less than the maximum number of potential sequences if each residue position was allowed to be any of the allowable 20 conventional amino acids (and/or allowable 5 unconventional amino/imino acids). A defined sequence kernal generally comprises variant and invariant residue positions and/or comprises variant residue positions which can comprise a residue selected from a defined subset of amino acid residues), and the like, either segmentally or over the entire length of the 10 individual selected library member sequence. Defined sequence kernals can refer to either amino acid sequences or polynucleotide sequences. For illustration and not limitation, the sequences $(NNK)_{10}$ and $(NNM)_{10}$, where N represents A, T, G, or C; K represents G or T; and M represents A or C, are defined 15 sequence kernals.

As used herein "epitope" refers to that portion of an antigen or other macromolecule capable of forming a binding interaction that interacts with the variable region binding pocket of an antibody. Typically, such binding interaction is 20 manifested as an intermolecular contact with one or more amino acid residues of a CDR.

As used herein, "receptor" refers to a molecule that has an affinity for a given ligand. Receptors can be naturally occurring or synthetic molecules. Receptors can be employed in 25 an unaltered state or as aggregates with other species. Receptors can be attached, covalently or noncovalently, to a binding member, either directly or via a specific binding substance. Examples of receptors include, but are not limited to, antibodies, including monoclonal antibodies and antisera 30 reactive with specific antigenic determinants (such as on viruses, cells, or other materials), cell membrane receptors, complex carbohydrates and glycoproteins, enzymes, and hormone receptors.

As used herein "ligand" refers to a molecule, such as 35 a random peptide or variable segment sequence, that is recognized by a particular receptor. As one of skill in the art will recognize, a molecule (or macromolecular complex) can be both a

receptor and a ligand. In general, the binding partner having a smaller molecular weight is referred to as the ligand and the binding partner having a greater molecular weight is referred to as a receptor.

5 As used herein, "linker" or "spacer" refers to a molecule or group of molecules that connects two molecules, such as a DNA binding protein and a random peptide, and serves to place the two molecules in a preferred configuration, e.g., so that the random peptide can bind to a receptor with minimal
10 steric hindrance from the DNA binding protein.

As used herein, the term "operably linked" refers to a linkage of polynucleotide elements in a functional relationship. A nucleic acid is "operably linked" when it is placed into a functional relationship with another nucleic acid
15 sequence. For instance, a promoter or enhancer is operably linked to a coding sequence if it affects the transcription of the coding sequence. Operably linked means that the DNA sequences being linked are typically contiguous and, where necessary to join two protein coding regions, contiguous and in
20 reading frame.

The term "recursive sequence recombination" as used herein refers to a method whereby a population of polynucleotide sequences are recombined with each other by any suitable recombination means (e.g., sexual PCR, homologous recombination,
25 site-specific recombination, etc.) to generate a library of sequence-recombined species which is then screened or subjected to selection to obtain those sequence-recombined species having a desired property; the selected species are then subjected to at least one additional cycle of recombination with themselves
30 and/or with other polynucleotide species and at subsequent selection or screening for the desired property.

Screening is, in general, a two-step process in which one first determines which cells do and do not express a screening marker and then physically separates the cells having
35 the desired property. Selection is a form of screening in which identification and physical separation are achieved simultaneously by expression of a selection marker, which, in

some genetic circumstances, allows cells expressing the marker to survive while other cells die (or vice versa). Screening markers include luciferase, β -galactosidase, and green fluorescent protein. Selection markers include drug and toxin resistance genes. Although spontaneous selection can and does occur in the course of natural evolution, in the present methods selection is performed by man.

Methodology

Nucleic acid shuffling is a method for recursive in vitro or in vivo homologous recombination of pools of nucleic acid fragments or polynucleotides. Mixtures of related nucleic acid sequences or polynucleotides are randomly or pseudorandomly fragmented, and reassembled to yield a library or mixed population of recombinant nucleic acid molecules or polynucleotides.

In contrast to cassette mutagenesis, only shuffling and error-prone PCR (or use of other mutation-enhancement methods; chemical mutagenesis, mutator strains, etc.) allow one to mutate a pool of sequences blindly (without sequence information other than primers).

The advantage of the mutagenic shuffling of this invention over error-prone PCR alone for repeated selection can best be explained with an example from antibody engineering. In Figure 1 is shown a schematic diagram of DNA shuffling as described herein. The initial library can consist of related sequences of diverse origin (i.e. antibodies from naive mRNA) or can be derived by any type of mutagenesis (including shuffling) of a single antibody gene. A collection of selected complementarity determining regions ("CDRs") is obtained after the first round of affinity selection (Fig.1). In the diagram the thick CDRs confer onto the antibody molecule increased affinity for the antigen. Shuffling allows the free combinatorial association of all of the CDR1s with all of the CDR2s with all of the CDR3s, etc. (Fig.1).

This method differs from PCR, in that it is an inverse chain reaction. In PCR, the number of molecules grows exponentially. In shuffling, however, the number of the polymerase start sites

and the number of molecules remains essentially the same. When dilution is used to allow further lengthening of the molecules, the shuffling process becomes an inverse chain reaction generating fewer molecules.

5 Since cross-overs occur at regions of homology, recombination will primarily occur between members of the same sequence family. This discourages combinations of CDRs that are grossly incompatible (eg. directed against different epitopes of the same antigen). It is contemplated that multiple families of
10 sequences can be shuffled in the same reaction. Further, shuffling conserves the relative order, such that, for example, CDR1 will not be found in the position of CDR2.

Rare shufflants will contain a large number of the best (eg. highest affinity) CDRs and these rare shufflants may be selected
15 based on their superior affinity (Fig. 1).

CDRs from a pool of 100 different selected antibody sequences can be permuted in up to 100^6 different ways. This large number of permutations cannot be represented in a single library of DNA sequences. Accordingly, it is contemplated that
20 multiple cycles of DNA shuffling and selection may be required depending on the length of the sequence and the sequence diversity desired.

Error-prone PCR, in contrast, keeps all the selected CDRs in the same relative sequence (Fig. 1), generating a much smaller
25 mutant cloud.

The template polynucleotide which may be used in the methods of this invention may be DNA or RNA. It may be of various lengths depending on the size of the gene or DNA fragment to be recombined or reassembled. Preferably the template
30 polynucleotide is from 50 bp to 50 kb. It is contemplated that entire vectors containing the nucleic acid encoding the protein of interest can be used in the methods of this invention, and in fact have been successfully used.

The template polynucleotide may be obtained by amplification
35 using the PCR reaction (U.S. Patent No. 4,683,202 and 4,683,195) or other amplification or cloning methods. However, the removal of free primers from the PCR product before fragmentation

provides a more efficient result. Failure to adequately remove the primers can lead to a low frequency of crossover clones.

The template polynucleotide often should be double-stranded. A double-stranded nucleic acid molecule is required to ensure 5 that regions of the resulting single-stranded nucleic acid fragments are complementary to each other and thus can hybridize to form a double-stranded molecule.

It is contemplated that single-stranded or double-stranded nucleic acid fragments having regions of identity to the template 10 polynucleotide and regions of heterology to the template polynucleotide may be added to the template polynucleotide at this step. It is also contemplated that two different but related polynucleotide templates can be mixed at this step.

The double-stranded polynucleotide template and any added 15 double-or single-stranded fragments are randomly digested into fragments of from about 5 bp to 5 kb or more. Preferably the size of the random fragments is from about 10 bp to 1000 bp, more preferably the size of the DNA fragments is from about 20 bp to 500 bp.

20 Alternatively, it is also contemplated that double-stranded nucleic acid having multiple nicks may be used in the methods of this invention. A nick is a break in one strand of the double-stranded nucleic acid. The distance between such nicks is preferably 5 bp to 5 kb, more preferably between 10 bp to 1000 25 bp.

The nucleic acid fragment may be digested by a number of different methods. The nucleic acid fragment may be digested with a nuclease, such as DNaseI or RNase. The nucleic acid may be randomly sheared by the method of sonication or by passage 30 through a tube having a small orifice.

It is also contemplated that the nucleic acid may also be partially digested with one or more restriction enzymes, such that certain points of cross-over may be retained statistically.

The concentration of any one specific nucleic acid fragment 35 will not be greater than 1% by weight of the total nucleic acid, more preferably the concentration of any one specific nucleic

acid sequence will not be greater than 0.1% by weight of the total nucleic acid.

The number of different specific nucleic acid fragments in the mixture will be at least about 100, preferably at least about 5 500, and more preferably at least about 1000.

At this step single-stranded or double-stranded nucleic acid fragments, either synthetic or natural, may be added to the random double-stranded nucleic acid fragments in order to increase the heterogeneity of the mixture of nucleic acid 10 fragments.

It is also contemplated that populations of double-stranded randomly broken or nicked nucleic acid fragments may be mixed or combined at this step. Damaged DNA can be exploited to enhance recombination via the nicked portions which can participate in 15 strand invasion, formation of recombination junctions, serve as free 3' ends for hybrid formation and the like.

Where insertion of mutations into the template polynucleotide is desired, single-stranded or double-stranded nucleic acid fragments having a region of identity to the 20 template polynucleotide and a region of heterology to the template polynucleotide may be added in a 20 fold excess by weight as compared to the total nucleic acid, more preferably the single-stranded nucleic acid fragments may be added in a 10 fold excess by weight as compared to the total nucleic acid.

Where a mixture of different but related template polynucleotides is desired, populations of nucleic acid fragments from each of the templates may be combined at a ratio of less than about 1:100, more preferably the ratio is less than about 1:40. For example, a backcross of the wild-type polynucleotide 30 with a population of mutated polynucleotide may be desired to eliminate neutral mutations (e.g., mutations yielding an insubstantial alteration in the phenotypic property being selected for). In such an example, the ratio of randomly digested wild-type polynucleotide fragments which may be added 35 to the randomly digested mutant polynucleotide fragments is approximately 1:1 to about 100:1, and more preferably from 1:1 to 40:1.

The mixed population of random length nucleic acid fragments are denatured to form single-stranded nucleic acid fragments and then reannealed. Only those single-stranded nucleic acid fragments having regions of homology with other single-stranded 5 nucleic acid fragments will reanneal.

The random length nucleic acid fragments may be denatured by heating. One skilled in the art could determine the conditions necessary to completely denature the double stranded nucleic acid. Preferably the temperature is from 80 °C to 100 °C, 10 more preferably the temperature is from 90 °C to 96 °C. Other methods which may be used to denature the nucleic acid fragments include pressure (36) and pH.

The nucleic acid fragments may be reannealed by cooling. Preferably the temperature is from 20 °C to 75 °C, more 15 preferably the temperature is from 40 °C to 65 °C. If a high frequency of crossovers is needed based on an average of only 4 consecutive bases of homology, recombination can be forced by using a low annealing temperature, although the process becomes more difficult. The degree of renaturation which occurs will 20 depend on the degree of homology between the population of single-stranded nucleic acid fragments.

Renaturation can be accelerated by the addition of polyethylene glycol ("PEG") or salt. The salt concentration is preferably from 0 mM to about 400 mM, more preferably the salt 25 concentration is from 10 mM to 100 mM. The salt may be KCl or NaCl. The concentration of PEG is preferably from 0% to 20%, more preferably from 5% to 10%. Higher concentrations of salt and/or PEG can be used, if desired.

The annealed nucleic acid fragments are next incubated in 30 the presence of a nucleic acid polymerase and dNTP's (i.e. dATP, dCTP, dGTP and dTTP). The nucleic acid polymerase may be the Klenow fragment, the Taq polymerase or any other DNA polymerase known in the art.

The approach to be used for the assembly depends on the 35 minimum degree of homology that should still yield crossovers. If the areas of identity are large, Taq polymerase can be used with an annealing temperature of between 45-65°C. If the areas

of identity are small, Klenow polymerase can be used with an annealing temperature of between 20-30°C. One skilled in the art could vary the temperature of annealing to increase the number of cross-overs achieved.

5 The polymerase may be added to the random nucleic acid fragments prior to annealing, simultaneously with annealing or after annealing.

The cycle of denaturation, renaturation and incubation in the presence of polymerase can be referred to as shuffling or 10 reassembly of the nucleic acid. This cycle is repeated for a desired number of times. Preferably the cycle is repeated from 2 to 50 times, more preferably the sequence is repeated from 10 to 40 times. The term "shuffling" encompasses a broader range of recursive recombination processes which can include, but are not 15 obligated to, PCR amplification or similar amplification methods; thus, shuffling can involve homologous recombination, site-specific recombination, chimera formation (e.g., Levichkin et al. op.cit), and the like, so long as used recursively (i.e., for more than one cycle of sequence recombination) with selection 20 and/or screening. Non-deterministic recombination, such as general homologous recombination can be used in combination with or in place of deterministic recombination, such as site-specific recombination where the sites of recombination are known and/or defined.

25 The resulting nucleic acid is a larger double-stranded polynucleotide of from about 50 bp to about 100 kb, preferably the larger polynucleotide is from 500 bp to 50 kb.

This larger polynucleotide fragment may contain a number of 30 copies of a nucleic acid fragment having the same size as the template polynucleotide in tandem. This concatemeric fragment is then digested into single copies of the template polynucleotide. The result will be a population of nucleic acid fragments of approximately the same size as the template polynucleotide. The population will be a mixed population where 35 single or double-stranded nucleic acid fragments having an area of identity and an area of heterology have been added to the template polynucleotide prior to shuffling. Alternatively, the

concatemer can be introduced (e.g., via electroporation, lipofection, or the like) directly without monomerization. For large sequences, it can be desirable to subdivide the large sequence into several subportions which are separately shuffled 5 with other substantially similar portions, and the pool of resultant shuffled subportions are then ligated, typically in original order, to generate a pool of shuffled large sequences which can then be used for transformation of a host cell, or the like.

10 These fragment are then cloned into the appropriate vector and the ligation mixture used to transform bacteria.

It is contemplated that the single nucleic acid fragments may be obtained from the larger concatemeric nucleic acid fragment by amplification of the single nucleic acid fragments 15 prior to cloning by a variety of methods including PCR (U.S. Patent No. 4,683,195 and 4,683,202) rather than by digestion of the concatemer. Alternatively, the concatemer can be introduced (e.g., via electroporation, lipofection, or the like) directly without monomerization.

20 The vector used for cloning is not critical provided that it will accept a DNA fragment of the desired size. If expression of the DNA fragment is desired, the cloning vehicle should further comprise transcription and translation signals next to the site of insertion of the DNA fragment to allow expression of 25 the DNA fragment in the host cell. Preferred vectors include the pUC series and the pBR series of plasmids.

The resulting bacterial population will include a number of recombinant DNA fragments having random mutations. This mixed 30 population may be tested to identify the desired recombinant nucleic acid fragment. The method of selection will depend on the DNA fragment desired.

For example, if a DNA fragment which encodes for a protein with increased binding efficiency to a ligand is desired, the proteins expressed by each of the DNA fragments in the population 35 or library may be tested for their ability to bind to the ligand by methods known in the art (i.e. panning, affinity chromatography). If a DNA fragment which encodes for a protein

with increased drug resistance is desired, the proteins expressed by each of the DNA fragments in the population or library may be tested for their ability to confer drug resistance to the host organism. One skilled in the art, given knowledge of the desired 5 protein, could readily test the population to identify DNA fragments which confer the desired properties onto the protein.

It is contemplated that one skilled in the art could use a phage display system in which fragments of the protein are expressed as fusion proteins on the phage surface (Pharmacia, 10 Milwaukee WI). The recombinant DNA molecules are cloned into the phage DNA at a site which results in the transcription of a fusion protein, a portion of which is encoded by the recombinant DNA molecule. The phage containing the recombinant nucleic acid molecule undergoes replication and transcription in the cell. 15 The leader sequence of the fusion protein directs the transport of the fusion protein to the tip of the phage particle. Thus the fusion protein which is partially encoded by the recombinant DNA molecule is displayed on the phage particle for detection and selection by the methods described above.

20 It is further contemplated that a number of cycles of nucleic acid shuffling may be conducted with nucleic acid fragments from a subpopulation of the first population, which subpopulation contains DNA encoding the desired recombinant protein. In this manner, proteins with even higher binding 25 affinities or enzymatic activity could be achieved.

It is also contemplated that a number of cycles of nucleic acid shuffling may be conducted with a mixture of wild-type nucleic acid fragments and a subpopulation of nucleic acid from the first or subsequent rounds of nucleic acid shuffling in order 30 to remove any silent mutations from the subpopulation.

Any source of nucleic acid, in purified form can be utilized as the starting nucleic acid. Thus the process may employ DNA or RNA including messenger RNA, which DNA or RNA may be single or double stranded. In addition, a DNA-RNA hybrid which contains 35 one strand of each may be utilized. The nucleic acid sequence may be of various lengths depending on the size of the nucleic acid sequence to be mutated. Preferably the specific nucleic

acid sequence is from 50 to 50000 base pairs. It is contemplated that entire vectors containing the nucleic acid encoding the protein of interest may be used in the methods of this invention.

The nucleic acid may be obtained from any source, for example, from plasmids such a pBR322, from cloned DNA or RNA or from natural DNA or RNA from any source including bacteria, yeast, viruses and higher organisms such as plants or animals. DNA or RNA may be extracted from blood or tissue material. The template polynucleotide may be obtained by amplification using the polynucleotide chain reaction (PCR) (U.S. Patent no. 4,683,202 and 4,683,195). Alternatively, the polynucleotide may be present in a vector present in a cell and sufficient nucleic acid may be obtained by culturing the cell and extracting the nucleic acid from the cell by methods known in the art.

Any specific nucleic acid sequence can be used to produce the population of mutants by the present process. It is only necessary that a small population of mutant sequences of the specific nucleic acid sequence exist or be created prior to the present process.

The initial small population of the specific nucleic acid sequences having mutations may be created by a number of different methods. Mutations may be created by error-prone PCR. Error-prone PCR uses low-fidelity polymerization conditions to introduce a low level of point mutations randomly over a long sequence. Alternatively, mutations can be introduced into the template polynucleotide by oligonucleotide-directed mutagenesis. In oligonucleotide-directed mutagenesis, a short sequence of the polynucleotide is removed from the polynucleotide using restriction enzyme digestion and is replaced with a synthetic polynucleotide in which various bases have been altered from the original sequence. The polynucleotide sequence can also be altered by chemical mutagenesis. Chemical mutagens include, for example, sodium bisulfite, nitrous acid, hydroxylamine, hydrazine or formic acid. Other agents which are analogues of nucleotide precursors include nitrosoguanidine, 5-bromouracil, 2-aminopurine, or acridine. Generally, these agents are added to the PCR reaction in place of the nucleotide precursor thereby

mutating the sequence. Intercalating agents such as proflavine, acriflavine, quinacrine and the like can also be used. Random mutagenesis of the polynucleotide sequence can also be achieved by irradiation with X-rays or ultraviolet light. Generally, 5 plasmid DNA or DNA fragments so mutagenized are introduced into *E. coli* and propagated as a pool or library of mutant plasmids.

Alternatively the small mixed population of specific nucleic acids may be found in nature in that they may consist of different alleles of the same gene or the same gene from 10 different related species (i.e., cognate genes). Alternatively, they may be related DNA sequences found within one species, for example, the immunoglobulin genes.

Once the mixed population of the specific nucleic acid sequences is generated, the polynucleotides can be used directly 15 or inserted into an appropriate cloning vector, using techniques well-known in the art.

The choice of vector depends on the size of the polynucleotide sequence and the host cell to be employed in the methods of this invention. The templates of this invention may 20 be plasmids, phages, cosmids, phagemids, viruses (e.g., retroviruses, parainfluenzavirus, herpesviruses, reoviruses, paramyxoviruses, and the like), or selected portions thereof (e.g., coat protein, spike glycoprotein, capsid protein). For example, cosmids, phagemids, YACs, and BACs are preferred where 25 the specific nucleic acid sequence to be mutated is larger because these vectors are able to stably propagate large nucleic acid fragments.

If the mixed population of the specific nucleic acid sequence is cloned into a vector it can be clonally amplified by 30 inserting each vector into a host cell and allowing the host cell to amplify the vector. This is referred to as clonal amplification because while the absolute number of nucleic acid sequences increases, the number of mutants does not increase.

35 Parallel PCR

In parallel PCR a large number of different PCR reactions occur in parallel in the same vessel, with the products

of one reaction priming the products of another reaction. As the PCR products prime each other, the average product size increases with the number of PCR cycles.

By using multiple primers in parallel, sequences in excess of 50 kb can be amplified. Whole genes and whole plasmids can be assembled in a single tube from synthetic oligonucleotides by parallel PCR. Sequences can be randomly mutagenized at various levels by random fragmentation and reassembly of the fragments by mutual priming. Site-specific mutations can be introduced into long sequences by random fragmentation of the template followed by reassembly of the fragments in the presence of mutagenic oligonucleotides. A particularly useful application of parallel PCR is called sexual PCR.

In sexual PCR, also called DNA shuffling, parallel PCR is used to perform *in vitro* recombination on a pool of DNA sequences. A mixture of related but not identical DNA sequences (typically PCR products, restriction fragments or whole plasmids) is randomly fragmented, for example by DNaseI treatment. These random fragments are then reassembled by parallel PCR. As the random fragments and their PCR products prime each other, the average size of the fragments increases with the number of PCR cycles. Recombination, or crossover, occurs by template switching, such as when a DNA fragment derived from one template primes on the homologous position of a related but different template. For example, sexual PCR can be used to construct libraries of chimaeras of genes from different species ('zoo libraries'). Sexual PCR is useful for *in vitro* evolution of DNA sequences. The libraries of new mutant combinations that are obtained by sexual PCR are selected for the best recombinant sequences at the DNA, RNA, protein or small molecule level. This process of recombination, selection and amplification is repeated for as many cycles as necessary to obtain a desired property or function.

Most versions of parallel PCR do not use primers. The DNA fragments, whether synthetic, obtained by random digestion, or by PCR with primers, serve as the template as well as the primers. Because the concentration of each different end

sequence in the reassembly reaction is very low, the formation of primer dimer is not observed, and if erroneous priming occurs, it can only grow at the same rate as the correctly annealed product. Parallel PCR requires many cycles of PCR because only 5 half of the annealed pairs have extendable overhangs and the concentration of 3' ends is low.

Utility

The DNA shuffling method of this invention can be performed 10 blindly on a pool of unknown sequences. By adding to the reassembly mixture oligonucleotides (with ends that are homologous to the sequences being reassembled) any sequence mixture can be incorporated at any specific position into another sequence mixture. Thus, it is contemplated that mixtures of 15 synthetic oligonucleotides, PCR fragments or even whole genes can be mixed into another sequence library at defined positions. The insertion of one sequence (mixture) is independent from the insertion of a sequence in another part of the template. Thus, the degree of recombination, the homology required, and the 20 diversity of the library can be independently and simultaneously varied along the length of the reassembled DNA.

This approach of mixing two genes may be useful for the humanization of antibodies from murine hybridomas. The approach of mixing two genes or inserting mutant sequences into genes may 25 be useful for any therapeutically used protein, for example, interleukin I, antibodies, tPA, growth hormone, etc. The approach may also be useful in any nucleic acid for example, promoters or introns or 3' untranslated region or 5' untranslated regions of genes to increase expression or alter specificity of expression 30 of proteins. The approach may also be used to mutate ribozymes or aptamers.

Shuffling requires the presence of homologous regions separating regions of diversity. If the sequences to be shuffled are not substantially identical, it is typically preferable to 35 employ intron-based shuffling and/or site-specific recombination. Scaffold-like protein structures may be particularly suitable for shuffling. The conserved scaffold determines the overall folding

by self-association, while displaying relatively unrestricted loops that mediate the specific binding. Examples of such scaffolds are the immunoglobulin beta-barrel, and the four-helix bundle (24). This shuffling can be used to create scaffold-like 5 proteins with various combinations of mutated sequences for binding.

In Vitro Shuffling

The equivalents of some standard genetic matings may also 10 be performed by shuffling *in vitro*. For example, a 'molecular backcross' can be performed by repeated mixing of the mutant's nucleic acid with the wild-type nucleic acid while selecting for 15 the mutations of interest. As in traditional breeding, this approach can be used to combine phenotypes from different sources 20 into a background of choice. It is useful, for example, for the removal of neutral mutations that affect unselected characteristics (i.e. immunogenicity). Thus it can be useful to determine which mutations in a protein are involved in the enhanced biological activity and which are not, an advantage 25 which cannot be achieved by error-prone mutagenesis or cassette mutagenesis methods.

Large, functional genes can be assembled correctly from a mixture of small random fragments. This reaction may be of use 25 for the reassembly of genes from the highly fragmented DNA of fossils (25). In addition random nucleic acid fragments from fossils may be combined with nucleic acid fragments from similar genes from related species.

It is also contemplated that the method of this invention can be used for the *in vitro* amplification of a whole genome from 30 a single cell as is needed for a variety of research and diagnostic applications. DNA amplification by PCR is in practice limited to a length of about 40 kb. Amplification of a whole genome such as that of *E. coli* (5,000 kb) by PCR would require 35 about 250 primers yielding 125 forty kb fragments. This approach is not practical due to the unavailability of sufficient sequence data. On the other hand, random digestion of the genome with DNaseI, followed by gel purification of small fragments will

provide a multitude of possible primers. Use of this mix of random small fragments as primers in a PCR reaction alone or with the whole genome as the template should result in an inverse chain reaction with the theoretical endpoint (assuming dilution 5 and additional PCR) of a single concatemer containing many copies of the genome.

100 fold amplification in the copy number and an average fragment size of greater than 50 kb may be obtained when only random fragments are used (see Example 2). It is thought that 10 the larger concatemer is generated by overlap of many smaller fragments. The quality of specific PCR products obtained using synthetic primers will be indistinguishable from the product obtained from unamplified DNA. It is expected that this approach will be useful for the mapping of genomes.

15 The polynucleotide to be shuffled can be fragmented randomly or non-randomly, at the discretion of the practitioner.

In Vivo Shuffling

In an embodiment of in vivo shuffling, the mixed population 20 of the specific nucleic acid sequence is introduced into bacterial (e.g., Archeabacteria) or eukaryotic cells under conditions such that at least two different nucleic acid sequences are present in each host cell. The fragments can be introduced into the host cells by a variety of different methods. 25 The host cells can be transformed with the fragments using methods known in the art, for example treatment with calcium chloride. If the fragments are inserted into a phage genome, the host cell can be transfected with the recombinant phage genome having the specific nucleic acid sequences. Alternatively, the 30 nucleic acid sequences can be introduced into the host cell using electroporation, natural competence, transduction, transfection, lipofection, biolistics, conjugation, and the like or other suitable method of introducing a polynucleotide sequence into a cell.

35 In general, in this embodiment, the specific nucleic acids sequences will be present in vectors which are capable of stably replicating the sequence in the host cell. In addition, it is

contemplated that the vectors will encode a marker gene such that host cells having the vector can be selected or screened. This ensures that the mutated specific nucleic acid sequence can be recovered after introduction into the host cell. However, it is
5 contemplated that the entire mixed population of the specific nucleic acid sequences need not be present on a vector sequence. Rather only a sufficient number of sequences need be cloned into vectors to ensure that after introduction of the fragments into the host cells each host cell contains two vector species having
10 at least one related-sequence nucleic acid sequence present therein. It is also contemplated that rather than having a subset of the population of the specific nucleic acids sequences cloned into vectors, this subset may be already stably integrated into the host cell.

15 It has been found that when two fragments which have regions of identity are inserted into the host cells, homologous recombination occurs between the two fragments. Such recombination between the two mutated specific nucleic acid sequences will result in the production of substantially all
20 combinations of all or most of the mutations (as limited by library size and propagation efficiency, etc.).

It has also been found that the frequency of recombination is increased if some of the mutated specific nucleic acid sequences are present on linear nucleic acid molecules.
25 Therefore, in a preferred embodiment, some of the specific nucleic acid sequences are present on linear nucleic acid fragments. In an embodiment, the nucleic acid molecules are single-stranded or substantially single-stranded, such as single-stranded phage genomes which may or may not comprise heterologous sequences. M13 phage is an example of a suitable ssDNA template,
30 and M13 has the advantage that the M13 virus can be shuffled in prokaryotic cells (e.g., E. coli), and then used to transfer DNA into mammalian cells.

After transformation, the host cell transformants are placed
35 under selection to identify those host cell transformants which contain mutated specific nucleic acid sequences having the qualities desired. For example, if increased resistance to a

particular drug is desired then the transformed host cells may be subjected to increased concentrations of the particular drug and those transformants producing mutated proteins able to confer increased drug resistance will be selected. If the enhanced 5 ability of a particular protein to bind to a receptor is desired, then expression of the protein can be induced from the transformants and the resulting protein assayed in a ligand binding assay by methods known in the art to identify that subset of the mutated population which shows enhanced binding to the 10 ligand. Alternatively, the protein can be expressed in another system to ensure proper processing.

Once a subset of the first recombined specific nucleic acid sequences (daughter sequences) having the desired characteristics are identified, they are then subject to a second round of 15 recombination.

In the second cycle of recombination, the recombined specific nucleic acid sequences may be mixed with the original mutated specific nucleic acid sequences (parent sequences) and the cycle repeated as described above. In this way a set of 20 second recombined specific nucleic acids sequences can be identified which have enhanced characteristics or encode for proteins having enhanced properties. This cycle can be repeated a number of times as desired.

It is also contemplated that in the second or subsequent 25 recombination cycle, a backcross can be performed. A molecular backcross can be performed by mixing the desired specific nucleic acid sequences with a large number of the wild-type sequence, such that at least one wild-type nucleic acid sequence and a mutated nucleic acid sequence are present in the same host cell 30 after transformation. Recombination with the wild-type specific nucleic acid sequence will eliminate those neutral or weakly contributory mutations that may affect unselected characteristics such as immunogenicity but not the selected characteristics.

In another embodiment of this invention, it is contemplated 35 that during the first round a subset of the specific nucleic acid sequences can be fragmented prior to introduction into the host cell. The size of the fragments must be large enough to contain

some regions of identity with the other sequences so as to homologously recombine with the other sequences. These fragments, ssDNA or dsDNA, can be coated with RecA in vitro to promote hybridization and/or integration into the host DNA. The size of the fragments will range from 0.03 kb to 100 kb more preferably from 0.2 kb to 10 kb. It is also contemplated that in subsequent rounds, all of the specific nucleic acid sequences other than the sequences selected from the previous round may be cleaved into fragments prior to introduction into the host cells.

"Cleavage" may be by nuclease digestion, PCR amplification (via partial extension or stuttering), or other suitable means for generating partial length polynucleotides of parent sequence(s).

Fragmentation of the sequences can be accomplished by a variety of methods known in the art. The sequences can be randomly fragmented or fragmented at specific sites in the nucleic acid sequence. Random fragments can be obtained by breaking the nucleic acid or exposing it to harsh physical treatment (e.g., shearing or irradiation) or harsh chemical agents (e.g., by free radicals; metal ions; acid treatment to depurinate and cleave). Random fragments can also be obtained, in the case of DNA by the use of DNase or like nuclease, or by other means as discussed herein. The sequences can be cleaved at specific sites by the use of restriction enzymes. The fragmented sequences can be single-stranded or double-stranded. If the sequences were originally single-stranded they can be denatured with heat, chemicals or enzymes prior to insertion into the host cell. The reaction conditions suitable for separating the strands of nucleic acid are well known in the art. Furthermore, partial PCR extension, PCR stuttering, and other related methods for producing partial length copies of a parental sequence can be used to effect "fragmentation", e.g., to obtain a hybrid product which contains segments derived from different parental sequences.

The steps of this process can be repeated indefinitely, being limited only by the number of possible mutants which can be achieved. After a certain number of cycles, all possible mutants will have been achieved and further cycles are redundant.

In an embodiment the same mutated template nucleic acid is repeatedly recombined and the resulting recombinants selected for the desired characteristic.

Therefore, the initial pool or population of mutated 5 template nucleic acid is cloned into a vector capable of replicating in bacteria such as *E. coli*. The particular vector is not essential, so long as it is capable of autonomous replication in *E. coli* or integration into a host chromosome. In a preferred embodiment, the vector is designed to allow the 10 expression and production of any protein encoded by the mutated specific nucleic acid linked to the vector. It is also preferred that the vector contain a gene encoding for a selectable marker.

The population of vectors containing the pool of mutated nucleic acid sequences is introduced into the *E. coli* host cells. 15 The vector nucleic acid sequences may be introduced by transformation, transfection or infection in the case of phage. The concentration of vectors used to transform the bacteria is such that a number of vectors is introduced into each cell. Once present in the cell, the efficiency of homologous recombination 20 is such that homologous recombination occurs between the various vectors. This results in the generation of mutants (daughters) having a combination of mutations which differ from the original parent mutated sequences.

The host cells are then replicated, typically clonally, and 25 selected for the marker gene present on the vector. Only those cells having a plasmid will grow under the selection.

The host cells which contain a vector are then tested for the presence of favorable mutations. Such testing may consist of placing the cells under selective pressure, for example, if 30 the gene to be selected is an improved drug resistance gene. If the vector allows expression of the protein encoded by the mutated nucleic acid sequence, then such selection may include allowing expression of the protein so encoded, isolation of the protein and testing of the protein to determine whether, for 35 example, it binds with increased efficiency to the ligand of interest.

characterized recA protein is from *E. coli*, in addition to the wild-type protein a number of mutant recA-like proteins have been identified (e.g., recA803). Further, many organisms have recA-like recombinases with strand-transfer activities (e.g., Fugisawa et al., (1985) Nucl. Acids Res. 13: 7473; Hsieh et al., (1986) Cell 44: 885; Hsieh et al., (1989) J. Biol. Chem. 264: 5089; Fishel et al., (1988) Proc. Natl. Acad. Sci. USA 85: 3683; Cassuto et al., (1987) Mol. Gen. Genet. 208: 10; Ganea et al., (1987) Mol. Cell Biol. 7: 3124; Moore et al., (1990) J. Biol. Chem. 19: 11108; Keene et al., (1984) Nucl. Acids Res. 12: 3057; Kimiec, (1984) Cold Spring Harbor Symp. 48:675; Kimeic, (1986) Cell 44: 545; Kolodner et al., (1987) Proc. Natl. Acad. Sci. USA 84 :5560; Sugino et al., (1985) Proc. Natl. Acad. Sci. USA 82: 3683; Halbrook et al., (1989) J. Biol. Chem. 264: 21403; Eisen et al., (1988) Proc. Natl. Acad. Sci. USA 85: 7481; McCarthy et al., (1988) Proc. Natl. Acad. Sci. USA 85: 5854; Lowenhaupt et al., (1989) J. Biol. Chem. 264: 20568, which are incorporated herein by reference. Examples of such recombinase proteins include, for example but not limitation: recA, recA803, uvsX, and other recA mutants and recA-like recombinases (Roca, A.I. (1990) Crit. Rev. Biochem. Molec. Biol. 25: 415), sep1 (Kolodner et al. (1987) Proc. Natl. Acad. Sci. (U.S.A.) 84: 5560; Tishkoff et al. Molec. Cell. Biol. 11: 2593), RuvC (Dunderdale et al. (1991) Nature 354: 506), DST2, KEM1, XRN1 (Dykstra et al. (1991) Molec. Cell. Biol. 11: 2583), STP α /DST1 (Clark et al. (1991) Molec. Cell. Biol. 11: 2576), HPP-1 (Moore et al. (1991) Proc. Natl. Acad. Sci. (U.S.A.) 88: 9067), other eukaryotic recombinases (Bishop et al. (1992) Cell 69: 439; Shinohara et al. (1992) Cell 69: 457); incorporated herein by reference. RecA may be purified from *E. coli* strains, such as *E. coli* strains JC12772 and JC15369 (available from A.J. Clark and M. Madiraju, University of California-Berkeley). These strains contain the recA coding sequences on a "runaway" replicating plasmid vector present at a high copy number per cell. The recA803 protein is a high-activity mutant of wild-type recA. The art teaches several examples of recombinase proteins, for example, from *Drosophila*, yeast, plant, human, and non-human mammalian cells, including

proteins with biological properties similar to recA (i.e., recA-like recombinases).

RecA protein is typically obtained from bacterial strains that overproduce the protein: wild-type *E. coli* recA protein and mutant recA803 protein may be purified from such strains. Alternatively, recA protein can also be purchased from, for example, Pharmacia (Piscataway, NJ).

RecA protein forms a nucleoprotein filament when it coats a single-stranded DNA. In this nucleoprotein filament, one monomer of recA protein is bound to about 3 nucleotides. This property of recA to coat single-stranded DNA is essentially sequence independent, although particular sequences favor initial loading of recA onto a polynucleotide (e.g., nucleation sequences). The nucleoprotein filament(s) can be formed on essentially any sequence-related polypeptide to be shuffled and can be formed in cells (e.g., bacterial, yeast, or mammalian cells), forming complexes with both single-stranded and double-stranded DNA.

Site-specific recombination can be used to accomplish recursive sequence recombination. Typically, sequences to be shuffled are flanked by one or more site-specific recombination sequences, such as for example a FLP recombination target site (FRT) often consisting of the two inverted 13 base repeats and an 8 bp spacer (O'Gorman et al. (1991) *Science* 251: 1351; Parsons et al. (1990) *J. Biol. Chem.* 265: 4527; Amin et al. (1991) *Mol. Cell. Biol.* 11: 4497, incorporated herein by reference). When FRT sequences are employed, the FLP recombinase is typically also employed, either *in vitro* or expressed in a host cell wherein the sequences to be recombined are introduced or are already present. Alternatives to the FLP/FRT system include, but are not limited to, the cre/lox system of phage P1 (Hoess and Abremski (1985) *J. Mol. Biol.* 181: 351), the γ/δ resolvase (Steitz et al. (1990) *Quarterly Rev. Biophys.* 23: 205), the attB/attP system of λ (Nunes-Duby et al. (1987) *Cell* 50: 779), and like site-specific recombination systems from bacteriophages λ , ϕ 80, P22, P2, P4, P1, and other like site-specific recombination systems selected by the practitioner. Guidance regarding the integrase family of

recombinases is found in Argos et al. (1986) EMBO J. 5: 433, incorporated herein by reference.

Exonuclease

5 In one aspect, the improved shuffling method includes the addition of at least one additive which is an enzyme having an exonuclease activity which is active at removing non-templated nucleotides introduced at 3' ends of product polynucleotides in shuffling amplification reactions catalyzed by a non-proofreading 10 polymerase. An examples of a suitable enzyme having an exonuclease activity includes but is not limited to Pfu polymerase. Examples of exonucleases are:

Bal31

Bacteriophage Lambda exonuclease

15 E. coli Exonuclease I

E. coli Exonuclease III

E. coli Exonuclease VII

Bacteriophage T7 gene 6.

20 Stuttering

In an aspect, the improved shuffling method comprises the modification wherein at least one cycle of amplification (i.e., extension with a polymerase) of reannealed fragmented library member polynucleotides is conducted under conditions 25 which produce a substantial fraction, typically at least 20 percent or more, of incompletely extended amplification products. The amplification products, including the incompletely extended amplification products are denatured and subjected to at least one additional cycle of reannealing and amplification. This 30 variation, wherein at least one cycle of reannealing and amplification provides a substantial fraction of incompletely extended products, is termed "stuttering" and in the subsequent amplification round the incompletely extended products reanneal to and prime extension on different sequence-related template 35 species.

In an aspect, the improved shuffling method comprises the modification wherein at least one cycle of amplification is

conducted using a collection of overlapping single-stranded DNA fragments of varying lengths corresponding to a first polynucleotide species or set of related-sequence polynucleotide species, wherein each overlapping fragment can each hybridize to
5 and prime polynucleotide chain extension from a second polynucleotide species serving as a template, thus forming sequence-recombined polynucleotides, wherein said sequence-recombined polynucleotides comprise a portion of at least one first polynucleotide species with an adjacent portion of the
10 second polynucleotide species which serves as a template. In a variation, the second polynucleotide species serving as a template contains uracil (i.e., a Kunkel-type template) and is substantially non-replicable in cells. This aspect of the invention can also comprise at least two recursive cycles of this
15 variation. In one embodiment, recursive cycles of shuffling using the method of Levitchkin et al. (1995) Mol. Biol. 29: 572, which produces partial extension PCR fragments is used to generate chimeras from a pool of parental sequences which are recursively shuffled.

20 In an aspect, the improved shuffling method comprises the modification wherein at least one cycle of amplification is conducted with an additive or polymerase in suitable conditions which promote template switching. In an embodiment where Tag polymerase is employed for amplification, addition of recA or
25 other polymerases enhances template switching. Template-switching can also be increased by increasing the DNA template concentration, among other means known by those skilled in the art.

30 In an embodiment of the general method, libraries of sequence-recombined polynucleotides are generated from sequence-related polynucleotides which are naturally-occurring genes or alleles of a gene. In this aspect, at least three naturally-occurring genes and/or alleles which comprise regions of at least 50 consecutive nucleotides which have at least 70 percent
35 sequence identity, preferably at least 90 percent sequence identity, are selected from a pool of gene sequences, such as by hybrid selection or via computerized sequence analysis using

sequence data from a database. The selected sequences are obtained as polynucleotides, either by cloning or via DNA synthesis, and shuffled by any of the various embodiments of the invention.

5 In an embodiment of the invention, the method comprises the further step of removing non-shuffled products (e.g., parental sequences) from sequence-recombined polynucleotides produced by any of the disclosed shuffling methods. Non-shuffled products can be removed or avoided by performing amplification
10 with: (1) a first PCR primer which hybridizes to a first parental polynucleotide species but does not substantially hybridize to a second parental polynucleotide species, and (2) a second PCR primer which hybridizes to a second parental polynucleotide species but does not substantially hybridize to
15 the first parental polynucleotide species, such that amplification occurs on from templates comprising the portion of the first parental sequence which hybridizes to the first PCR primer and also comprising the portion of the second parental sequence which hybridizes to the second PCR primer, thus only
20 sequence-recombined polynucleotides are amplified.

In an embodiment of the invention, "bridging" genes can be synthesized. If two or more parental polynucleotides (e.g., genes) lack satisfactory sequence similarity for efficient homologous recombination or for efficient cross-priming for PCR
25 amplification, intermediate (or "bridging") genes can be synthesized which share sufficient sequence identity with the parental sequences. The bridging gene need not be active or confer a phenotype or selectable property, it need only provide a template having sufficient sequence identity to accomplish
30 shuffling of the parental sequences. The intermediate homology of the bridging gene, and the necessary sequence(s) can be determined by computer or manually.

The invention also provides additional formats for
35 performing recursive recombination *in vivo*, either in prokaryotic or eucaryotic cells. These formats include recombination between plasmids, recombination between viruses, recombination between

plasmid and virus, recombination between a chromosome and plasmid or virus and intramolecular recombination (e.g., between two sequences on a plasmid). Recursive recombination can be performed entirely *in vivo* whereby successive rounds of *in vivo* recombination are interspersed by rounds of selection or screening. *In vivo* formats can also be used in combination with *in vitro* formats. For example, one can perform one round of *in vitro* shuffling, a round of selection, a round of *in vivo* shuffling, a further round of selection, a further round of *in vitro* shuffling and a further round of selection and so forth. The various *in vivo* formats are now considered in turn.

(a) Plasmid-Plasmid Recombination

The initial substrates for recombination are a collection of polynucleotides comprising variant forms of a gene. The variant forms usually show substantial sequence identity to each other sufficient to allow homologous recombination between substrates. The diversity between the polynucleotides can be natural (e.g., allelic or species variants), induced (e.g., error-prone PCR, synthetic genes, codon-usage altered sequence variants), or the result of *in vitro* recombination. There should be at least sufficient diversity between substrates that recombination can generate more diverse products than there are starting materials. There must be at least two substrates differing in at least two positions. However, commonly a library of substrates of 10^3 - 10^8 members is employed. The degree of diversity depends on the length of the substrate being recombined and the extent of the functional change to be evolved. Diversity at between 0.1-25% of positions is typical.

The diverse substrates are incorporated into plasmids. The plasmids are often standard cloning vectors, e.g., bacterial multicopy plasmids. However, in some methods to be described below, the plasmids include MOB functions. The substrates can be incorporated into the same or different plasmids. Often at least two different types of plasmid having different types of selection marker are used to allow selection for cells containing at least two types of vector. Also, where different types of

plasmid are employed, the different plasmids can come from two distinct incompatibility groups to allow stable co-existence of two different plasmids within the cell. Nevertheless, plasmids from the same incompatibility group can still co-exist within the 5 same cell for sufficient time to allow homologous recombination to occur.

Plasmids containing diverse substrates are initially introduced into cells by any transfection methods (e.g., chemical transformation, natural competence, transduction, electroporation 10 or biolistics). Often, the plasmids are present at or near saturating concentration (with respect to maximum transfection capacity) to increase the probability of more than one plasmid entering the same cell.

The plasmids containing the various substrates can be transfected 15 simultaneously or in multiple rounds. For example, in the latter approach cells can be transfected with a first aliquot of plasmid, transfectants selected and propagated, and then infected with a second aliquot of plasmid.

Having introduced the plasmids into cells, recombination 20 between substrates to generate recombinant genes occurs within cells containing multiple different plasmids merely by propagating the cells. However, cells that receive only one plasmid are less able to participate in recombination and the potential contribution of substrates on such plasmids to 25 evolution is wasted. The rate of evolution can be increased by allowing all substrates to participate in recombination. Such can be achieved by subjecting transfected cells to electroporation. The conditions for electroporation are the same 30 as those conventionally used for introducing exogenous DNA into cells (e.g., 1,000-2,500 volts, 400 µF and a 1-2 mM gap). Under these conditions, plasmids are exchanged between cells allowing all substrates to participate in recombination. In addition the products of recombination can undergo further rounds of recombination with each other or with the original substrate. 35 The rate of evolution can also be increased by use of conjugative transfer. To exploit conjugative transfer, substrates can be cloned into plasmids having MOB genes and tra genes are also

provided in cis or in trans to the MOB genes. The effect of conjugative transfer is very similar to electroporation in that it allows plasmids to move between cells and allows recombination between any substrate, and the products of previous recombination 5 to occur merely by propagating the culture. The details of how conjugative transfer is exploited in these vectors are discussed in more detail below. The rate of evolution can also be increased by use of mutator host cells (e.g., Mut L, S, D, T, H; human ataxia telangiectasia cells).

10 The time for which cells are propagated and recombination is allowed to occur, of course, varies with the cell type but is generally not critical, because even a small degree of recombination can substantially increase diversity relative to the starting materials. Cells bearing plasmids containing 15 recombinant genes are subject to screening or selection for a desired function. For example, if the substrate being evolved contains a drug resistance gene, one would select for drug resistance. Cells surviving screening or selection can be subjected to one or more rounds of screening/selection followed 20 by recombination or can be subjected directly to an additional round of recombination.

The next round of recombination can be achieved by several different formats independently of the previous round. For example, a further round of recombination can be effected simply 25 by resuming the electroporation or conjugation-mediated intercellular transfer of plasmids described above. Alternatively, a fresh substrate or substrates, the same or different from previous substrates, can be transfected into cells surviving selection/screening. Optionally, the new substrates 30 are included in plasmid vectors bearing a different selective marker and/or from a different incompatibility group than the original plasmids. As a further alternative, cells surviving selection/screening can be subdivided into two subpopulations, plasmid DNA extracted from one subpopulation and transfected into 35 the other, where the substrates from the plasmids from the two subpopulations undergo a further round of recombination. In either of the latter two options, the rate of evolution can be

increased by employing electroporation, conjugation or mutator cells, as described above. In a still further variation, DNA from cells surviving screening/selection can be extracted and subjected to *in vitro* DNA shuffling.

5 After the second round of recombination, a second round of screening/selection is performed, preferably under conditions of increased stringency. If desired, further rounds of recombination and selection/screening can be performed using the same strategy as for the second round. With successive rounds
10 of recombination and selection/screening, the surviving recombined substrates evolve toward acquisition of a desired phenotype. Typically, in this and other methods of *in vivo* recursive recombination, the final product of recombination that has acquired the desired phenotype differs from starting
15 substrates at 0.1%-25% of positions and has evolved at a rate orders of magnitude in excess (e.g., by at least 10-fold, 100-fold, 1000-fold, or 10,000 fold) of the rate of naturally acquired mutation of about 1 mutation per 10^{-9} positions per generation (see Anderson & Hughes, Proc. Natl. Acad. Sci. USA 93,
20 906-907 (1996)).

Fig. 26 shows an exemplary scheme of plasmid-plasmid recombination. Panel A of the figure shows a library of variant genes cloned into a plasmid. The library is then introduced into cells. Some cells take up a single plasmid and other cells take
25 up two plasmids as shown in panel B. For cells having taken up two plasmids, the plasmids recombine to give the products shown in panel C. Plasmids can then be transferred between cells by electroporation or conjugation as shown in panel D, and further recombination can occur to give the products shown in panel E.
30 Screening/selection then isolates plasmids bearing genes that have evolved toward acquisition of the property that selection/screening is designed to identify. In the course of selection, a cell bearing two plasmids of which only one contributes to the selected phenotype, may lose the other
35 plasmid, as shown in panel F.

(b) Virus-Plasmid Recombination

The strategy used for plasmid-plasmid recombination can also be used for virus-plasmid recombination; usually, phage-plasmid recombination. However, some additional comments particular to the use of viruses are appropriate. The initial substrates for 5 recombination are cloned into both plasmid and viral vectors. It is usually not critical which substrate(s) are inserted into the viral vector and which into the plasmid, although usually the viral vector should contain different substrate(s) from the plasmid. As before, the plasmid typically contains a selective 10 marker. The plasmid and viral vectors can both be introduced into cells by transfection as described above. However, a more efficient procedure is to transfet the cells with plasmid, select transfecants and infect the transfecants with virus. Because the efficiency of infection of many viruses approaches 15 100% of cells, most cells transfected and infected by this route contain both a plasmid and virus bearing different substrates.

Homologous recombination occurs between plasmid and virus generating both recombined plasmids and recombined virus. For 20 some viruses, such as filamentous phage, in which intracellular DNA exists in both double-stranded and single-stranded forms, both can participate in recombination. Provided that the virus is not one that rapidly kills cells, recombination can be augmented by use of electroporation or conjugation to transfer 25 plasmids between cells. Recombination can also be augmented for some types of virus by allowing the progeny virus from one cell to reinfect other cells. For some types of virus, virus infected-cells show resistance to superinfection. However, such resistance can be overcome by infecting at high multiplicity 30 and/or using mutant strains of the virus in which resistance to superinfection is reduced.

The result of infecting plasmid-containing cells with virus depends on the nature of the virus. Some viruses, such as filamentous phage, stably exist with a plasmid in the cell and 35 also extrude progeny phage from the cell. Other viruses, such as lambda having a cosmid genome, stably exist in a cell like plasmids without producing progeny virions. Other viruses, such

as the T-phage and lytic lambda, undergo recombination with the plasmid but ultimately kill the host cell and destroy plasmid DNA. For viruses that infect cells without killing the host, cells containing recombinant plasmids and virus can be 5 screened/selected using the same approach as for plasmid-plasmid recombination. Progeny virus extruded by cells surviving selection/screening can also be collected and used as substrates in subsequent rounds of recombination. For viruses that kill their host cells, recombinant genes resulting from recombination 10 reside only in the progeny virus. If the screening or selective assay requires expression of recombinant genes in a cell, the recombinant genes should be transferred from the progeny virus to another vector, e.g., a plasmid vector, and retransfected into cells before selection/screening is performed.

15 For filamentous phage, the products of recombination are present in both cells surviving recombination and in phage extruded from these cells. The dual source of recombinant products provides some additional options relative to the plasmid-plasmid recombination. For example, DNA can be isolated 20 from phage particles for use in a round of *in vitro* recombination. Alternatively, the progeny phage can be used to transfect or infect cells surviving a previous round of screening/selection, or fresh cells transfected with fresh substrates for recombination. In an aspect, the invention 25 employs recombination between multiple single-stranded species, such as single-stranded bacteriophages and/or phagemids.

Fig. 27 illustrates a scheme for virus-plasmid recombination. Panel A shows a library of variant forms of gene cloned into plasmid and viral vectors. The plasmids are then 30 introduced into cells as shown in panel B. The viral genomes are packaged *in vitro* and used to infect the cells in panel B. The viral genomes can undergo replication within the cell, as shown in panel C. The viral genomes undergo recombination with plasmid genomes generating the plasmid and viral forms shown in panel D. 35 Both plasmids and viral genomes can undergo further rounds of replication and recombination generating the structures shown in panels E and F. Screening/selection identifies cells containing

plasmid and/or viral genomes having genes that have evolved best to allow survival of the cell in the screening/selection process, as shown in panel G. These viral genomes are also present in viruses extruded by such cells.

5

(c) Virus-Virus Recombination

The principles described for plasmid-plasmid and plasmid-viral recombination can be applied to virus-virus recombination with a few modifications. The initial substrates for 10 recombination are cloned into a viral vector. Usually, the same vector is used for all substrates. Preferably, the virus is one that, naturally or as a result of mutation, does not kill cells. After insertion, viral genomes are usually packaged in vitro. The packaged viruses are used to infect cells at high 15 multiplicity such that there is a high probability that a cell will receive multiple viruses bearing different substrates.

After the initial round of infection, subsequent steps depend on the nature of infection as discussed in the previous section.

20 For example, if the viruses have phagemid genomes such as lambda cosmids or M13, F1 or Fd phagemids, the phagemids behave as plasmids within the cell and undergo recombination simply by propagating the cells. Recombination is particularly efficient between single-stranded forms of intracellular DNA. 25 Recombination can be augmented by electroporation of cells. Following selection/screening, cosmids containing recombinant genes can be recovered from surviving cells (e.g., by heat induction of a cos⁻ lysogenic host cell), repackaged in vitro, and used to infect fresh cells at high multiplicity for a further 30 round of recombination.

If the viruses are filamentous phage, recombination of replicating form DNA occurs by propagating the culture of infected cells. Selection/screening identifies colonies of cells containing viral vectors having recombinant genes with improved 35 properties, together with phage extruded from such cells. Subsequent options are essentially the same as for plasmid-viral recombination.

Fig. 28 shows an example of virus-virus recombination. A library of diverse genes is cloned into a lambda cosmid. The recombinant cosmid DNA is packaged in vitro and used to infect host cells at high multiplicity such that many cosmids bearing 5 different inserts enter the same cell. The cell chosen is a cos-lambda lysogen, which on induction, packages cosmid DNA without packaging lysogenic DNA. Cosmids recombine within the cell. Recombination can be accelerated by the use of host cells that are MutD, MutS, MutL and/or express a modified recA. Induction 10 of the lysogen results in release of packaged recombinant cosmids, having greater diversity than the starting materials.

(d) Chromosome-Plasmid Recombination

This format can be used to evolve both the chromosomal and 15 plasmid-borne substrates. The format is particularly useful in situations in which many chromosomal genes contribute to a phenotype or one does not know the exact location of the chromosomal gene(s) to be evolved. The initial substrates for recombination are cloned into a plasmid vector. If the 20 chromosomal gene(s) to be evolved are known, the substrates constitute a family of sequences showing a high degree of sequence identity but some divergence from the chromosomal gene. If the chromosomal genes to be evolved have not been located, the 25 initial substrates usually constitute a library of DNA segments of which only a small number show sequence identity to the gene or gene(s) to be evolved. Divergence between plasmid-borne substrate and the chromosomal gene(s) can be induced by mutagenesis or by obtaining the plasmid-borne substrates from a different species than that of the cells bearing the chromosome.

30 The plasmids bearing substrates for recombination are transfected into cells having chromosomal gene(s) to be evolved. Evolution can occur simply by propagating the culture, and can be accelerated by transferring plasmids between cells by 35 conjugation or electroporation. Evolution can be further accelerated by use of mutator host cells or by seeding a culture of nonmutator host cells being evolved with mutator host cells

and inducing intercellular transfer of plasmids by electroporation or conjugation. Preferably, mutator host cells used for seeding contain a negative selection marker to facilitate isolation of a pure culture of the nonmutator cells being evolved. Selection/screening identifies cells bearing chromosomes and/or plasmids that have evolved toward acquisition of a desired function.

Subsequent rounds of recombination and selection/screening proceed in similar fashion to those described for plasmid-plasmid recombination. For example, further recombination can be effected by propagating cells surviving recombination in combination with electroporation or conjugative transfer of plasmids. Alternatively, plasmids bearing additional substrates for recombination can be introduced into the surviving cells. Preferably, such plasmids are from a different incompatibility group and bear a different selective marker than the original plasmids to allow selection for cells containing at least two different plasmids. As a further alternative, plasmid and/or chromosomal DNA can be isolated from a subpopulation of surviving cells and transfected into a second subpopulation. Chromosomal DNA can be cloned into a plasmid vector before transfection.

Fig. 29 illustrates a scheme for plasmid-chromosome shuffling. Panel A shows variant forms of a gene cloned into a plasmid vector. The plasmids are introduced into cells as shown in panel B. In the cells, the plasmids replicate and undergo recombination with a chromosomal copy of the gene, as shown in panel C. Exchange of plasmids between cells can be effected by electroporation or conjugation as shown in panel D. The chromosomal genes in the two cells shown in panel D have evolved to different variant forms. Screening/selection identifies the cell bearing the chromosomal gene that has evolved that has acquired a desired property that allows the cell to survive screening/selection, as shown in panel E.

As in the other methods described above, the virus is usually one that does not kill the cells, and is often a phage or phagemid. The procedure is substantially the same as for plasmid-chromosome recombination. Substrates for recombination 5 are cloned into the vector. Vectors including the substrates can then be transfected into cells or in vitro packaged and introduced into cells by infection. Viral genomes recombine with host chromosomes merely by propagating a culture. Evolution can be accelerated by allowing intercellular transfer of viral 10 genomes by electroporation, or reinfection of cells by progeny virions. Screening/selection identifies cells having chromosomes and/or viral genomes that have evolved toward acquisition of a desired function.

There are several options for subsequent rounds of 15 recombination. For example, viral genomes can be transferred between cells surviving selection/recombination by electroporation. Alternatively, viruses extruded from cells surviving selection/screening can be pooled and used to superinfect the cells at high multiplicity. Alternatively, fresh 20 substrates for recombination can be introduced into the cells, either on plasmid or viral vectors.

e. Evolution of Genes by Conjugative Transfer

As noted above, the rate of in vivo evolution of plasmids 25 DNA can be accelerated by allowing transfer of plasmids between cells by conjugation. Conjugation is the transfer of DNA occurring during contact between cells. See Guiney (1993) in: *Bacterial Conjugation* (Clewell, ed., Plenum Press, New York), pp. 30 75-104; Reimann & Haas in *Bacterial Conjugation* (Clewell, ed., Plenum Press, New York 1993), at pp.137-188 (incorporated by reference in their entirety for all purposes). Conjugation occurs between many types of gram negative bacteria, and some 35 types of gram positive bacteria. Conjugative transfer is also known between bacteria and plant cells (*Agrobacterium tumefaciens*) or yeast.

As discussed in copending application attorney docket no. 16528J-014612, the genes responsible for conjugative transfer can themselves be evolved to expand the range of cell types (e.g., from bacteria to mammals) between which such transfer can occur.

Conjugative transfer is effected by an origin of transfer (oriT) and flanking genes (MOB A, B and C), and 15-25 genes, termed tra, encoding the structures and enzymes necessary for conjugation to occur. The transfer origin is defined as the site required in cis for DNA transfer. Tra genes include tra A, B, C, D, E, F, G, H, I, J, K, L, M, N, P, Q, R, S, T, U, V, W, X, Y, Z, vir AB (alleles 1-11), C, D, E, G, IHF, and FinOP. OriT is sometimes also designated as a tra gene. Other cellular enzymes, including those of the RecBCD pathway, RecA, SSB protein, DNA gyrase, DNA polI, and DNA ligase, are also involved in conjugative transfer. RecE or recF pathways can substitute for RecBCD.

The tra genes and MOB genes can be expressed in cis or trans to oriT. Vectors undergoing conjugation also have an origin of replication which is classified as belonging to an incompatibility group such as Inc A, B, C, D, E, F (I-VI), H (I, Y), i (1, 2, 5, ALPHA), J, K, L, M, N, P (ALPHA, BETA, 1 ALPHA, 3, 7, 10, 13) Q, R (H1, H2, H3) S, T, U, W, X, Z. Only vectors from different incompatibility groups can stably co-exist in the same cell. However, when two vectors from the same incompatibility group are transfected into the same cell, the vectors transiently coexist for sufficient time that recombination can occur between the vectors.

One structural protein encoded by a tra gene is the sex pilus, a filament constructed of an aggregate of a single polypeptide protruding from the cell surface. The sex pilus binds to a polysaccharide on recipient cells and forms a conjugative bridge through which DNA can transfer. This process activates a site-specific nuclease encoded by a MOB gene, which specifically cleaves DNA to be transferred at oriT. The cleaved DNA is then threaded through the conjugation bridge by the action of other tra enzymes.

DNA is transferred more efficiently between cells when present as a component of the mobilizable vector. However, some mobilizable vectors integrate into the host chromosome and thereby mobilize adjacent genes from the chromosome. The F plasmid of *E. coli*, for example, integrates into the chromosome at high frequency and mobilizes genes unidirectional from the site of integration. Other mobilizable vectors do not spontaneously integrate into a host chromosome at high efficiency but can be induced to do by growth under particular conditions (e.g., treatment with a mutagenic agent, growth at a nonpermissive temperature for plasmid replication). See Reimann & Haas in *Bacterial Conjugation* (ed. Clewell, Plenum Press, NY 1993), Ch. 6.

Conjugation provides a means of recombining gene(s) *in vivo* to generate diverse recombinant forms of the gene(s). As in other methods of recursive recombination, iterative cycles of recombination and selection/screening can be used to evolve the gene(s) toward acquisition of a new or improved property. As in any method of recursive recombination, the first step is to generate a library of diverse forms of the gene or genes to be evolved. The diverse forms can be the result of natural diversity, the application of traditional mutagenesis methods (e.g., error-prone PCR or cassette mutagenesis) or the result of any of the other recombination formats discussed in this application, or any combination of these. The number of diverse forms can vary widely from about 10 to 100, 10^4 , 10^6 , 10^8 or 10^{10} . Often, the gene(s) of interest are mutagenized as discrete units. However, if the location of gene(s) is not known or a large number of genes are to be evolved simultaneously, initial diversity can be generated by *in situ* mutagenesis of a chromosome containing the gene(s).

The library of diverse forms of a gene or gene(s) is introduced into cells containing the apparatus necessary for conjugative transfer (assuming that the library is not already contained in such cells), usually in an arrangement such that the genes can be expressed. For example, if the gene(s) are mutagenized in the absence of essential regulatory sequences such

as promoter, these sequences are reattached before introduction into cells. Similarly, if a fragment of a gene is mutagenized in isolation, the mutagenesis products are usually reassociated with unchanged flanking sequences before being introduced into 5 cells. The apparatus necessary for conjugative transfer comprises a vector having an origin of transfer together with the mob and tra genes whose expression is necessary for conjugative transfer to occur. These genes can be included in the vector, in one or more different vectors, or in the chromosome. The 10 library of diverse forms of the gene to be evolved is usually inserted into the vector containing the origin of transfer (see Fig. 30). However, in some situations the library of diverse forms of the gene can be present in the chromosome or a second vector, as well as, or instead of in the vector containing the 15 origin of transfer. The library of diverse forms can be inserted in different places in different cells.

A vector bearing a library of variant forms contains at least one origin of replication. If transfer between different cell types is contemplated, the vector can contain two origins 20 of replication, one functional in each cell type (i.e., a shuttle vector). Alternatively, if it is intended that transferred genes should integrate into the chromosome of recipient cells, it is preferable that the vector not contain an origin of replication functional in the recipient cells (i.e., a suicide vector). The 25 oriT site and/or MOB genes can be introduced into a vector by cloning or transposing the RK2/RP4 MOB function (Guiney, *J. Mol. Biol.* 162, 699-703 (1982)), or by cointegrate formation with a MOB-containing plasmid. A convenient method for large plasmids is to use 'Tn5-Mob', which is the Tn5 transposon containing the 30 oriT of RP4. For example, pUC-like mobilizable vectors pK18 and pK19 (Schafer et al. (1995) *Gene* 145:69-73) are suitable starting vectors for cloning the tra gene library to be evolved.

Although not necessary, recombination is sometimes facilitated by inserting the diverse gene library into two 35 different kinds of vectors having different incompatibility origins. Each vector should have an oriT and the cell should express MOB and tra functions suitable for mobilization of both

vectors. Use of two such kinds of vectors allows stable coexistence of multiple vectors within the same cell and increases the efficiency of recombination between the vectors.

The collection of cells is propagated in any suitable media 5 to allow gene expression to occur. Tra and mob genes are expressed and mediate transfer of the mobilizable vector between cells. If the diverse library is cloned into a mobilizable vector, its members are transferred as components of the vector. If the diverse library, or certain elements of the library, are 10 in trans to the mobilizable vector, they are transferred only if the mobilizable vector integrates into or proximate to the elements. As discussed above, integration frequently occurs spontaneously for the *E. coli* F plasmid and can be induced for other mobilizable vectors.

As a result of transfer of members of the diverse library 15 between cells, some of the cells come to contain more than one member of the diverse library. The multiple members undergo recombination within such cells generating still further diversity in a library of recombinant forms. In general, the longer cells are propagated the more recombinant forms are 20 generated. Generally, recombination results in more than one recombination product within the same cell. If both recombination products are on vectors and the vectors are from the same incompatibility group, one of the vectors is lost as the 25 cells are propagated. This process occurs faster if the cells are propagated on selective media in which one or other of the recombinant products confers a selective advantage. After a suitable period of recombination, which depends on the cell type 30 and its growth cycle time, the recombinant forms are subject to screening or selection. Because the recombinant forms are already present in cells, this format for recombination is particularly amenable to alternation with cycles of *in vivo* screening or selection. The conditions for screening or 35 selection, of course, depend on the property which it is desired that the gene(s) being evolved acquire or improve in. For example, if the property is drug resistance, recombinant forms having the best drug resistance can be selected by exposure to

the drug. Alternatively, if a cluster of genes is being evolved to produce a drug as a secondary metabolite, cells bearing recombinant clusters of the genes can be screened by overlaying colonies of cell bearing recombinant cluster with a lawn of cells
5 that are sensitive to the drug. Colonies having recombinant clusters resulting in production of the best drug are identified from holes in the lawn. If the gene being evolved confers enhanced growth characteristics, cells bearing the best genes can be selected by growth competition. Antibiotic production can be
10 a growth rate advantage if cells are competing with other cell types for growth.

Screening/selection produces a subpopulation of cells expressing recombinant forms of gene(s) that have evolved toward acquisition of a desired property. These recombinant forms can
15 then be subjected to further rounds of recombination and screening/selection in any order. For example, a second round of screening/selection can be performed analogous to the first resulting in greater enrichment for genes having evolved toward acquisition of the desired property. Optionally, the stringency
20 of selection can be increased between rounds (e.g., if selecting for drug resistance, the concentration of drug in the media can be increased). Further rounds of recombination can also be performed by an analogous strategy to the first round generating further recombinant forms of the gene(s). Alternatively, further
25 rounds of recombination can be performed by any of the other molecular breeding formats discussed. Eventually, a recombinant form of the gene(s) is generated that has fully acquired the desired property.

Fig. 30 provides an example of how a drug resistance gene
30 can be evolved by conjugative transfer. Panel A shows a library of diverse genes cloned into a mobilizable vector bearing as oriT. The vectors are present in cells containing a second vector which provides tra functions. Conjugative transfer results in movement of the mobilizable vectors between cells,
35 such that different vectors bearing different variant forms of a gene occupy the same cell, as shown in panel B. The different forms of the gene recombine to give the products shown in panel

C. After conjugation and recombination has proceeded for a desired time, cells are selected to identify those containing the recombinant genes, as shown in panel D.

In one aspect, the alternative shuffling method 5 includes the use of intra-plasmidic recombination, wherein libraries of sequence-recombined polynucleotide sequences are obtained by genetic recombination *in vivo* of direct sequence repeats located on the same plasmid. In a variation, the sequences to be recombined are flanked by site-specific 10 recombination sequences and the polynucleotides are present in a site-specific recombination system, such as an integron (Hall and Collins (1995) Mol. Microbiol. **15**: 593, incorporated herein by reference).

In an aspect of the invention, mutator strains of host 15 cells are used to enhance recombination of more highly mismatched sequence-related polynucleotides. Bacterial strains such as MutL, MutS, or MutH or other cells expressing the Mut proteins (XL-1red; Stratagene, San Diego, CA) can be used as host cells for shuffling of sequence-related polynucleotides by *in vivo* 20 recombination. Other mutation-prone host cell types can also be used, such as those having a proofreading-defective polymerase (Foster et al. (1995) Proc. Natl. Acad. Sci. (U.S.A.) **92**: 7951, incorporated herein by reference). Other *in vivo* mutagenic 25 formats can be employed, including administering chemical or radiological mutagens to host cells. Examples of such mutagens include but are not limited to: ENU, MMNG, nitrosourea, BuDR, and the like.

Shuffling can be used to evolve polymerases capable of incorporation of base analogs in PCR or PCR-like amplification 30 reactions. A DNA polymerase which is evolved to use base analogs can be used to copy DNA by PCR into a chemical form which gives more resolvable fragmentation patterns in mass spectrometry, such as for mass spectrometry DNA sequencing. The base analogs can have fewer and/or more favorable fragmentation sites to enhance 35 or facilitate the interpretation of the mass spectrum patterns.

Variant polymerases can also be evolved by recursive sequence recombination to incorporate non-natural nucleotides or

nucleotide analogs, such as phosphorothioate nucleotides. Phosphorothioate nucleotides made with such variant polymerases can provide many uses, including naked DNA gene therapy vectors which are resistant to nuclease degradation. Other examples of 5 properties of polymerases which can be modified via recursive sequence recombination include, but are not limited to, processivity, error rate, proofreading, thermal stability, oxidation resistance, nucleotide preferences, template specificity, and the like, among others.

10 In an embodiment, fluorescence-activated cell sorting or analogous methodology is used to screen for host cells, typically mammalian cells, insect cells, or bacterial cells, comprising a library member of a recursively recombined sequence library, wherein the host cell having a library member conferring 15 a desired phenotype can be selected on the basis of fluorescence or optical density at one or more detection wavelengths. In one embodiment, for example, each library member typically encodes an enzyme, which may be secreted from the cell or may be intracellular, and the enzyme catalyzes conversion of a 20 chromogenic or fluorogenic substrate, which may be capable of diffusing into the host cell (e.g., if said enzyme is not secreted). Host cells containing library members are contained in fluid drops or gel drops and passed by a detection apparatus where the drops are illuminated with an excitation wavelength and 25 a detector measures either fluorescent emission wavelength radiation and/or measures optical density (absorption) at one or more excitatory wavelength(s). The cells suspended in drops are passed across a sample detector under conditions wherein only about one individual cell is present in a sample detection zone 30 at a time. A source, illuminates each cell and a detector, typically a photomultiplier or photodiode, detects emitted radiation. The detector controls gating of the cell in the detection zone into one of a plurality of sample collection regions on the basis of the signal(s) detected. A general 35 description of FACS apparatus and methods is provided in U.S. Patents 4,172,227; 4,347,935; 4,661,913; 4,667,830; 5,093,234; 5,094,940; and 5,144,224, incorporated herein by reference . A

suitable alternative to conventional FACS is available from One Cell Systems, Inc. Cambridge, MA.

As can be appreciated from the disclosure above, the present invention has a wide variety of applications. Accordingly, the
5 following examples are offered by way of illustration, not by way of limitation.

EXPERIMENTAL EXAMPLES

In the examples below, the following abbreviations have the following meanings. If not defined below, then the abbreviations have their art recognized meanings.

5	ml	=	milliliter
	μ l	=	microliters
	μ M	=	micromolar
	nM	=	nanomolar
	PBS	=	phosphate buffered saline
10	ng	=	nanograms
	μ g	=	micrograms
	IPTG	=	isopropylthio- β -D-galactoside
	bp	=	basepairs
	kb	=	kilobasepairs
15	dNTP	=	deoxynucleoside triphosphates
	PCR	=	polymerase chain reaction
	X-gal	=	5-bromo-4-chloro-3-indolyl- β -D-galactoside
	DNAseI	=	deoxyribonuclease
	PBS	=	phosphate buffered saline
20	CDR	=	complementarity determining regions
	MIC	=	minimum inhibitory concentration
	scFv	=	single-chain Fv fragment of an antibody

In general, standard techniques of recombination DNA technology are described in various publications, e.g. Sambrook et al., 1989, Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory; Ausubel et al., 1987, Current Protocols in Molecular Biology, vols. 1 and 2 and supplements, and Berger and Kimmel, Methods in Enzymology, Volume 152, Guide to Molecular Cloning Techniques (1987), Academic Press, Inc., San Diego, CA, each of which is incorporated herein in their entirety by reference. Restriction enzymes and polynucleotide modifying enzymes were used according to the manufacturers recommendations. Oligonucleotides were synthesized on an Applied Biosystems Inc. Model 394 DNA synthesizer using ABI chemicals. If desired, PCR amplimers for amplifying a predetermined DNA sequence may be selected at the discretion of the practitioner.

EXAMPLES40 Example 1. LacZ alpha gene reassembly

1) Substrate preparation

The substrate for the reassembly reaction was the dsDNA polymerase chain reaction ("PCR") product of the wild-type LacZ

alpha gene from pUC18. (Fig. 2) (28; Gene Bank No. X02514) The primer sequences were 5'AAAGCGTCGATTTTGAT3' (SEQ ID NO:1) and 5'ATGGGGTTCCGCACATTT3' (SEQ ID NO:2). The free primers were removed from the PCR product by Wizard PCR prep (Promega, Madison WI) according to the manufacturer's directions. The removal of the free primers was found to be important.

5 2) DNaseI digestion

About 5 μ g of the DNA substrate was digested with 0.15 units of DNaseI (Sigma, St. Louis MO) in 100 μ l of [50 mM Tris-HCl pH 10 7.4, 1 mM MgCl₂], for 10-20 minutes at room temperature. The digested DNA was run on a 2% low melting point agarose gel. Fragments of 10-70 basepairs (bp) were purified from the 2% low melting point agarose gels by electrophoresis onto DE81 ion exchange paper (Whatman, Hillsborough OR). The DNA fragments 15 were eluted from the paper with 1 M NaCl and ethanol precipitated.

15 3) DNA Reassembly

The purified fragments were resuspended at a concentration of 10 - 30 ng/ μ l in PCR Mix (0.2 mM each dNTP, 2.2 mM MgCl₂, 50 20 mM KCl, 10 mM Tris-HCl pH 9.0, 0.1% Triton X-100, 0.3 μ l Taq DNA polymerase, 50 μ l total volume). No primers were added at this 25 point. A reassembly program of 94°C for 60 seconds, 30-45 cycles of [94°C for 30 seconds, 50-55°C for 30 seconds, 72°C for 30 seconds] and 5 minutes at 72°C was used in an MJ Research (Watertown MA) PTC-150 thermocycler. The PCR reassembly of small fragments into larger sequences was followed by taking samples of the reaction after 25, 30, 35 ,40 and 45 cycles of reassembly (Fig. 2).

Whereas the reassembly of 100-200 bp fragments can yield a 30 single PCR product of the correct size, 10-50 base fragments typically yield some product of the correct size, as well as products of heterogeneous molecular weights. Most of this size heterogeneity appears to be due to single-stranded sequences at the ends of the products, since after restriction enzyme 35 digestion a single band of the correct size is obtained.

4) PCR with primers

After dilution of the reassembly product into the PCR Mix with 0.8 μ M of each of the above primers (SEQ ID Nos: 1 and 2) and about 15 cycles of PCR, each cycle consisting of [94°C for 5 30 seconds, 50°C for 30 seconds and 72°C for 30 seconds], a single product of the correct size was obtained (Fig. 2).

5) Cloning and analysis

The PCR product from step 4 above was digested with the terminal restriction enzymes BamHI and EcoO109 and gel purified 10 as described above in step 2. The reassembled fragments were ligated into pUC18 digested with BamHI and EcoO109. *E. coli* were transformed with the ligation mixture under standard conditions as recommended by the manufacturer (Stratagene, San Diego CA) and plated on agar plates having 100 μ g/ml ampicillin, 0.004% X-gal 15 and 2mM IPTG. The resulting colonies having the *Hin*DIII-*Nhe*I fragment which is diagnostic for the ++ recombinant were identified because they appeared blue.

This Example illustrates that a 1.0 kb sequence carrying the LacZ alpha gene can be digested into 10-70 bp fragments, and that 20 these gel purified 10-70 bp fragments can be reassembled to a single product of the correct size, such that 84% (N=377) of the resulting colonies are LacZ⁺ (versus 94% without shuffling; Fig. 2).

The DNA encoding the LacZ gene from the resulting LacZ⁻ 25 colonies was sequenced with a sequencing kit (United States Biochemical Co., Cleveland OH) according to the manufacturer's instructions and the genes were found to have point mutations due to the reassembly process (Table 1). 11/12 types of substitutions were found, and no frameshifts.

TABLE 1
Mutations introduced by mutagenic shuffling

Transitions	Frequency	Transversions	Frequency
G - A	6	A - T	1
5 A - G	4	A - C	2
	7	C - A	1
	3	C - G	0
		G - C	3
10 T - C		G - T	2
		T - A	1
		T - G	2

A total of 4,437 bases of shuffled lacZ DNA were sequenced.

15 The rate of point mutagenesis during DNA reassembly from 10-70 bp pieces was determined from DNA sequencing to be 0.7% (N=4,473), which is similar to error-prone PCR. Without being limited to any theory it is believed that the rate of point 20 mutagenesis may be lower if larger fragments are used for the reassembly, or if a proofreading polymerase is added.

When plasmid DNA from 14 of these point-mutated LacZ- colonies were combined and again reassembled/shuffled by the method described above, 34% (N=291) of the resulting colonies 25 were LacZ+, and these colonies presumably arose by recombination of the DNA from different colonies.

The expected rate of reversal of a single point mutation by error-prone PCR, assuming a mutagenesis rate of 0.7% (10), would be expected to be <1%.

30 Thus large DNA sequences can be reassembled from a random mixture of small fragments by a reaction that is surprisingly efficient and simple. One application of this technique is the recombination or shuffling of related sequences based on homology.

35 Example 2. LacZ gene and whole plasmid DNA shuffling

1) LacZ gene shuffling

Crossover between two markers separated by 75 bases was measured using two LacZ gene constructs. Stop codons were 40 inserted in two separate areas of the LacZ alpha gene to serve as negative markers. Each marker is a 25 bp non-homologous sequence with four stop codons, of which two are in the LacZ gene

reading frame. The 25 bp non-homologous sequence is indicated in Figure 3 by a large box. The stop codons are either boxed or underlined. A 1:1 mixture of the two 1.0 kb LacZ templates containing the +- and -+ versions of the LacZ alpha gene (Fig. 5) was digested with DNaseI and 100-200 bp fragments were purified as described in Example 1. The shuffling program was conducted under conditions similar to those described for reassembly in Example 1 except 0.5 μ l of polymerase was added and the total volume was 100 μ l.

10 After cloning, the number of blue colonies obtained was 24%; (N=386) which is close to the theoretical maximum number of blue colonies (i.e. 25%), indicating that recombination between the two markers was complete. All of the 10 blue colonies contained the expected *HindIII-NheI* restriction fragment.

15 2) Whole plasmid DNA shuffling

Whole 2.7 kb plasmids (pUC18-+ and pUC18+-) were also tested. A 1:1 mixture of the two 2.9 kb plasmids containing the +- and -+ versions of the LacZ alpha gene (Fig. 3) was digested with DNaseI and 100-200 bp fragments were purified as described 20 in Example 1. The shuffling program was conducted under conditions similar to those described for reassembly in step (1) above except the program was for 60 cycles [94°C for 30 seconds, 55°C for 30 seconds, 72°C for 30 seconds]. Gel analysis showed that after the shuffling program most of the product was greater 25 than 20 kb. Thus, whole 2.7 kb plasmids (pUC18 -- and pUC18 +-) were efficiently reassembled from random 100-200 bp fragments without added primers.

After digestion with a restriction enzyme having a unique site on the plasmid (*EcoO109*), most of the product consisted of 30 a single band of the expected size. This band was gel purified, religated and the DNA used to transform *E. coli*. The transformants were plated on 0.004% X-gal plates as described in Example 1. 11% (N= 328) of the resulting plasmids were blue and thus ++ recombinants.

35 3) Spiked DNA Shuffling

Oligonucleotides that are mixed into the shuffling mixture can be incorporated into the final product based on the homology

of the flanking sequences of the oligonucleotide to the template DNA (Fig. 4). The LacZ⁻ stop codon mutant (pUC18 -+) described above was used as the DNaseI digested template. A 66 mer oligonucleotide, including 18 bases of homology to the wild-type 5 LacZ gene at both ends was added into the reaction at a 4-fold molar excess to correct stop codon mutations present in the original gene. The shuffling reaction was conducted under conditions similar to those in step 2 above. The resulting product was digested, ligated and inserted into *E. coli* as 10 described above.

Table 2

	% blue colonies
15 Control	0.0 (N>1000)
Top strand spike	8.0 (N=855)
Bottom strand spike	9.3 (N=620)
Top and bottom strand spike	2.1 (N=537)

20 ssDNA appeared to be more efficient than dsDNA, presumably due to competitive hybridization. The degree of incorporation can be varied over a wide range by adjusting the molar excess, annealing temperature, or the length of homology.

25 Example 3. DNA reassembly in the complete absence of primers

Plasmid pUC18 was digested with restriction enzymes EcoRI, EcoO109, XmnI and AlwNI, yielding fragments of approximately 370, 460, 770 and 1080 bp. These fragments were electrophoresed and 30 separately purified from a 2% low melting point agarose gel (the 370 and 460 basepair bands could not be separated), yielding a large fragment, a medium fragment and a mixture of two small fragments in 3 separate tubes.

Each fragment was digested with DNaseI as described in 35 Example 1, and fragments of 50-130 bp were purified from a 2% low melting point agarose gel for each of the original fragments.

PCR mix (as described in Example 1 above) was added to the purified digested fragments to a final concentration of 10 ng/ μ l of fragments. No primers were added. A reassembly reaction was 40 performed for 75 cycles [94°C for 30 seconds, 60°C for 30 seconds] separately on each of the three digested DNA fragment

mixtures, and the products were analyzed by agarose gel electrophoresis.

The results clearly showed that the 1080, 770 and the 370 and 460 bp bands reformed efficiently from the purified 5 fragments, demonstrating that shuffling does not require the use of any primers at all.

Example 4. IL-1 β gene shuffling

This example illustrates that crossovers based on homologies 10 of less than 15 bases may be obtained. As an example, a human and a murine IL-1 β gene were shuffled.

A murine IL1- β gene (BBG49) and a human IL1- β gene with *E. coli* codon usage (BBG2; R&D Systems, Inc., Minneapolis MN) were used as templates in the shuffling reaction. The areas of 15 complete homology between the human and the murine IL-1 β sequences are on average only 4.1 bases long (Fig. 5, regions of heterology are boxed).

Preparation of dsDNA PCR products for each of the genes, removal of primers, DNaseI digestion and purification of 10-50 20 bp fragments was similar to that described above in Example 1. The sequences of the primers used in the PCR reaction were 5'TTAGGCACCCAGGCTTT3' (SEQ ID NO:3) and 5'ATGTGCTGCAAGGCGATT3' (SEQ ID NO:4).

The first 15 cycles of the shuffling reaction were performed 25 with the Klenow fragment of DNA polymerase I, adding 1 unit of fresh enzyme at each cycle. The DNA was added to the PCR mix of Example 1 which mix lacked the polymerase. The manual program was 94°C for 1 minute, and then 15 cycles of: [95°C for 1 minute, 30 10 seconds on dry ice/ethanol (until frozen), incubate about 20 seconds at 25°C , add 1U of Klenow fragment and incubate at 25°C for 2 minutes]. In each cycle after the denaturation step, the tube was rapidly cooled in dry ice/ethanol and reheated to the annealing temperature. Then the heat-labile polymerase was added. The enzyme needs to be added at every cycle. Using this 35 approach, a high level of crossovers was obtained, based on only a few bases of uninterrupted homology (Fig. 5, positions of cross-overs indicated by " _ | ").

After these 15 manual cycles, Taq polymerase was added and an additional 22 cycles of the shuffling reaction [94°C for 30 seconds, 35°C for 30 seconds] without primers were performed.

The reaction was then diluted 20-fold. The following 5 primers were added to a final concentration of 0.8 μM: 5' AACGCCGCATGCAAGCTTGGATCCTTATT3' (SEQ ID NO:5) and 5' AAAGCCCTCTAGATGATTACGAATTCAAT3' (SEQ ID NO:6) and a PCR reaction was performed as described above in Example 1. The second primer pair differed from the first pair only because a 10 change in restriction sites was deemed necessary.

After digestion of the PCR product with *Xba*I and *Sph*I, the fragments were ligated into *Xba*I-*Sph*I-digested pUC18. The sequences of the inserts from several colonies were determined by a dideoxy DNA sequencing kit (United States Biochemical Co., 15 Cleveland OH) according to the manufacturer's instructions.

A total of 17 crossovers were found by DNA sequencing of nine colonies. Some of the crossovers were based on only 1-2 bases of uninterrupted homology.

It was found that to force efficient crossovers based on 20 short homologies, a very low effective annealing temperature is required. With any heat-stable polymerase, the cooling time of the PCR machine (94°C to 25°C at 1-2 degrees/second) causes the effective annealing temperature to be higher than the set annealing temperature. Thus, none of the protocols based on Tag 25 polymerase yielded crossovers, even when a ten-fold excess of one of the IL1-β genes was used. In contrast, a heat-labile polymerase, such as the Klenow fragment of DNA polymerase I, can be used to accurately obtain a low annealing temperature.

30 Example 5. DNA shuffling of the TEM-1 betalactamase gene

The utility of mutagenic DNA shuffling for directed molecular evolution was tested in a betalactamase model system. TEM-1 betalactamase is a very efficient enzyme, limited in its reaction rate primarily by diffusion. This example determines 35 whether it is possible to change its reaction specificity and obtain resistance to the drug cefotaxime that it normally does not hydrolyze.

The minimum inhibitory concentration (MIC) of cefotaxime on bacterial cells lacking a plasmid was determined by plating 10 µl of a 10⁻² dilution of an overnight bacterial culture (about 1000 cfu) of *E. coli* XL1-blue cells (Stratagene, San Diego CA) 5 on plates with varying levels of cefotaxime (Sigma, St. Louis MO), followed by incubation for 24 hours at 37°C.

Growth on cefotaxime is sensitive to the density of cells, and therefore similar numbers of cells needed to be plated on each plate (obtained by plating on plain LB plates). Platings 10 of 1000 cells were consistently performed.

1) Initial Plasmid Construction

A pUC18 derivative carrying the bacterial TEM-1 betalactamase gene was used (28). The TEM-1 betalactamase gene confers resistance to bacteria against approximately 0.02 µg/ml 15 of cefotaxime. *Sfi*I restriction sites were added 5' of the promoter and 3' of the end of the gene by PCR of the vector sequence with two primers:

Primer A (SEQ ID NO: 7) :
5' TTCTATTGACGGCCTGTCAGGCCTCATATATAACTTTAGATTGATTT3' and Primer B
20 (SEQ ID NO: 8) :
5' TTGACGGCACTGGCCATGGTGGCCAAAAATAAACAAATAGGGTTCCGCGCACATT3'
and by PCR of the betalactamase gene sequence with two other primers:

Primer C (SEQ ID NO:9) :
25 5' AACTGACCACGGCCTGACAGGCCGGTCTGACAGTTACCAATGCTT, and
Primer D (SEQ ID NO:10) :
5' AACCTGTCCTGGCCACCATGGCCTAAATACATTCAAATATGTTAT.

The two reaction products were digested with *Sfi*I, mixed, ligated and used to transform bacteria.

30 The resulting plasmid was pUC182Sfi. This plasmid contains an *Sfi*I fragment carrying the TEM-1 gene and the P-3 promoter.

The minimum inhibitory concentration of cefotaxime for *E. coli* XL1-blue (Stratagene, San Diego CA) carrying this plasmid 35 was 0.02 µg/ml after 24 hours at 37°C.

The ability to improve the resistance of the betalactamase gene to cefotaxime without shuffling was determined by stepwise

replating of a diluted pool of cells (approximately 10^7 cfu) on 2-fold increasing drug levels. Resistance up to 1.28 $\mu\text{g}/\text{ml}$ could be obtained without shuffling. This represented a 64 fold increase in resistance.

5 2) DNaseI digestion

The substrate for the first shuffling reaction was dsDNA of 0.9 kb obtained by PCR of pUC182Sfi with primers C and D, both of which contain a *Sfi*I site.

10 The free primers from the PCR product were removed by Wizard PCR prep (Promega, Madison WI) at every cycle.

About 5 μg of the DNA substrate(s) was digested with 0.15 units of DNaseI (Sigma, St. Louis MO) in 100 μl of 50 mM Tris-HCl pH 7.4, 1 mM MgCl₂, for 10 min at room temperature. Fragments of 100-300 bp were purified from 2% low melting point agarose 15 gels by electrophoresis onto DE81 ion exchange paper (Whatman, Hillsborough OR), elution with 1 M NaCl and ethanol precipitation by the method described in Example 1.

3) Gene shuffling

The purified fragments were resuspended in PCR mix (0.2 mM 20 each dNTP, 2.2 mM MgCl₂, 50 mM KCl, 10 mM Tris-HCl pH 9.0, 0.1% Triton X-100), at a concentration of 10 - 30 ng/ μl . No primers were added at this point. A reassembly program of 94°C for 60 seconds, then 40 cycles of [94°C for 30 seconds, 50-55°C for 30 25 seconds, 72°C for 30 seconds] and then 72°C for 5 minutes was used in an MJ Research (Watertown MA) PTC-150 thermocycler.

4) Amplification of Reassembly Product with primers

After dilution of the reassembly product into the PCR mix with 0.8 μM of each primer (C and D) and 20 PCR cycles [94°C for 30 seconds, 50°C for 30 seconds, 72°C for 30 seconds] a single 30 product 900 bp in size was obtained.

5) Cloning and analysis

After digestion of the 900 bp product with the terminal restriction enzyme *Sfi*I and agarose gel purification, the 900 bp product was ligated into the vector pUC182Sfi at the unique *Sfi*I 35 site with T4 DNA ligase (BRL, Gaithersburg MD). The mixture was electroporated into *E. coli* XL1-blue cells and plated on LB plates with 0.32-0.64 $\mu\text{g}/\text{ml}$ of cefotaxime (Sigma, St. Louis MO).

The cells were grown for up to 24 hours at 37°C and the resulting colonies were scraped off the plate as a pool and used as the PCR template for the next round of shuffling.

6) Subsequent Reassembly Rounds

5 The transformants obtained after each of three rounds of shuffling were plated on increasing levels of cefotaxime. The colonies (>100, to maintain diversity) from the plate with the highest level of cefotaxime were pooled and used as the template for the PCR reaction for the next round.

10 A mixture of the cefotaxime^r colonies obtained at 0.32-0.64 µg/ml in Step (5) above were used as the template for the next round of shuffling. 10 ul of cells in LB broth were used as the template in a reassembly program of 10 minutes at 99°C, then 35 cycles of [94°C for 30 seconds, 52°C for 30 seconds, 72°C for 30
15 seconds] and then 5 minutes at 72°C as described above.

The reassembly products were digested and ligated into pUC182Sfi as described in step (5) above. The mixture was electroporated into *E. coli* XL1-blue cells and plated on LB plates having 5-10 µg/ml of cefotaxime.

20 Colonies obtained at 5-10 µg/ml were used for a third round similar to the first and second rounds except the cells were plated on LB plates having 80-160 µg/ml of cefotaxime. After the third round, colonies were obtained at 80-160 µg/ml, and after replating on increasing concentrations of cefotaxime, colonies
25 could be obtained at up to 320 µg/ml after 24 hours at 37°C (MIC=320 µg/ml).

Growth on cefotaxime is dependent on the cell density, requiring that all the MICs be standardized (in our case to about 1,000 cells per plate). At higher cell densities, growth at up
30 to 1280 µg/ml was obtained. The 5 largest colonies grown at 1,280 µg/ml were plated for single colonies twice, and the Sfil inserts were analyzed by restriction mapping of the colony PCR products.

One mutant was obtained with a 16,000 fold increased
35 resistance to cefotaxime (MIC=0.02 µg/ml to MIC=320 µg/ml).

After selection, the plasmid of selected clones was transferred back into wild-type *E. coli* XL1-blue cells

(Stratagene, San Diego CA) to ensure that none of the measured drug resistance was due to chromosomal mutations.

Three cycles of shuffling and selection yielded a 1.6×10^4 -fold increase in the minimum inhibitory concentration of the extended broad spectrum antibiotic cefotaxime for the TEM-1 betalactamase. In contrast, repeated plating without shuffling resulted in only a 16-fold increase in resistance (error-prone PCR or cassette mutagenesis).

7) Sequence analysis

All 5 of the largest colonies grown at 1,280 $\mu\text{g}/\text{ml}$ had a restriction map identical to the wild-type TEM-1 enzyme. The *Sfi*I insert of the plasmid obtained from one of these colonies was sequenced by dideoxy DNA sequencing (United States Biochemical Co., Cleveland OH) according to the manufacturer's instructions. All the base numbers correspond to the revised pBR322 sequence (29), and the amino acid numbers correspond to the ABL standard numbering scheme (30). The amino acids are designated by their three letter codes and the nucleotides by their one letter codes. The term G4205A means that nucleotide 4205 was changed from guanidine to adenine.

Nine single base substitutions were found. G4205A is located between the -35 and -10 sites of the betalactamase P3 promoter (31). The promoter up-mutant observed by Chen and Clowes (31) is located outside of the *Sfi*I fragment used here, and thus could not have been detected. Four mutations were silent (A3689G, G3713A, G3934A and T3959A), and four resulted in an amino acid change (C3448T resulting in Gly238Ser, A3615G resulting in Met182Thr, C3850T resulting in Glu104Lys, and G4107A resulting in Ala18Val).

30 8) Molecular Backcross

Molecular backcrossing with an excess of the wild-type DNA was then used in order to eliminate non-essential mutations.

Molecular backcrossing was conducted on a selected plasmid from the third round of DNA shuffling by the method identical to normal shuffling as described above, except that the DNaseI digestion and shuffling reaction were performed in the presence of a 40-fold excess of wild-type TEM-1 gene fragment. To make

the backcross more efficient, very small DNA fragments (30 to 100 bp) were used in the shuffling reaction. The backcrossed mutants were again selected on LB plates with 80-160 µg/ml of cefotaxime (Sigma, St. Louis MO).

5 This backcross shuffling was repeated with DNA from colonies from the first backcross round in the presence of a - fold excess of wild-type TEM-1 DNA. Small DNA fragments (30- 0 bp) were used to increase the efficiency of the backcross. The second round of backcrossed mutants were again selected on LB
10 plates with 80-160 µg/ml of cefotaxime.

The resulting transformants were plated on 160 µg/ml of cefotaxime, and a pool of colonies was replated on increasing levels of cefotaxime up to 1,280 µg/ml. The largest colony obtained at 1,280 µg/ml was replated for single colonies.

15 This backcrossed mutant was 32,000 fold more resistant than wild-type. (MIC=640 µg/ml) The mutant strain is 64-fold more resistant to cefotaxime than previously reported clinical or engineered TEM-1-derived strains. Thus, it appears that DNA shuffling is a fast and powerful tool for at least several cycles
20 of directed molecular evolution.

The DNA sequence of the SfiI insert of the backcrossed mutant was determined using a dideoxy DNA sequencing kit (United States Biochemical Co., Cleveland OH) according to the manufacturer's instructions (Table 3). The mutant had 9 single
25 base pair mutations. As expected, all four of the previously identified silent mutations were lost, reverting to the sequence of the wild-type gene. The promoter mutation (G4205A) as well as three of the four amino acid mutations (Glu104Lys, Met182Thr, and Gly238Ser) remained in the backcrossed clone, suggesting that
30 they are essential for high level cefotaxime resistance. However, two new silent mutations (T3842C and A3767G), as well as three new mutations resulting in amino acid changes were found (C3441T resulting in Arg241His, C3886T resulting in Gly92Ser, and G4035C resulting in Ala42Gly). While these two silent mutations
35 do not affect the protein primary sequence, they may influence prot in expr ssion level (for example by mRNA structure) and possibly even protein folding (by changing the codon usage and

therefore the pause site, which has been implicated in protein folding).

Table 3
Mutations in Betalactamase

Mutation Type	Non-Backcrossed	Backcrossed
5 amino acid change	Ala18Lys Glu104Lys Met182Thr Gly238Ser	- Glu104Lys Met182Thr Gly238Ser
10	- -	Ala42Gly Gly92Ser
15 silent	T3959A G3934A G3713A A3689G	- - - -
20 promoter	G4205A	G4205A

Both the backcrossed and the non-backcrossed mutants have a promoter mutation (which by itself or in combination results in a 2-3 fold increase in expression level) as well as three common amino acid changes (Glu104Lys, Met182Thr and Gly238Ser). Glu104Lys and Gly238Ser are mutations that are present in several cefotaxime resistant or other TEM-1 derivatives (Table 4).

9) Expression Level Comparison

The expression level of the betalactamase gene in the wild-type plasmid, the non-backcrossed mutant and in the backcrossed mutant was compared by SDS-polyacrylamide gel electrophoresis (4-20%; Novex, San Diego CA) of periplasmic extracts prepared by osmotic shock according to the method of Witholt, B. (32).

Purified TEM-1 betalactamase (Sigma, St. Louis MO) was used as a molecular weight standard, and *E. coli* XL1-blue cells lacking a plasmid were used as a negative control.

The mutant and the backcrossed mutant appeared to produce a 2-3 fold higher level of the betalactamase protein compared to the wild-type gene. The promoter mutation appeared to result in a 2-3 times increase in betalactamase.

Example 6. Construction of mutant combinations of the TEM-1 betalactamase gene

To determine the resistance of different combinations of mutations and to compare the new mutants to published mutants, several mutants were constructed into an identical plasmid background. Two of the mutations, Glu104Lys and Gly238Ser, are known as cefotaxime mutants. All mutant combinations constructed had the promoter mutation, to allow comparison to selected mutants. The results are shown in Table 4.

Specific combinations of mutations were introduced into the wild-type pUC182Sfi by PCR, using two oligonucleotides per mutation.

The oligonucleotides to obtain the following mutations were:

Ala42Gly

(SEQ ID NO:11) AGTTGGGTGGACGAGTGGGTTACATCGAACT and (SEQ ID NO:12)
15 AACCCACTCGTCCACCCAACTGATCTTCAGCAT;

Gln39Lys:

(SEQ ID NO:13) AGTAAAAGATGCTGAAGATAAGTTGGGTGCAC GAGTGGGTT and
(SEQ ID NO:14) ACTTATCTTCAGCATCTTTACTT;

Gly92Ser:

20 (SEQ ID NO:15) AAGAGCAACTCAGTGCCGCATACACTATTCT and (SEQ ID NO:16) ATGGCGGCGACTGAGTTGCTCTTGCCCCGGCGTCAAT;

Glu104Lys:

(SEQ ID NO:17) TATTCTCAGAATGACTTGGTTAAGTACTCACCAGT CACAGAA and
(SEQ ID NO:18) TTAACCAAGTCATTCTGAGAAT;

25 **Met182Thr:**

(SEQ ID NO:19) AACGACGAGCGTGACACCACGACGCCTGTAGCAATG and (SEQ ID NO:20) TCGTGGTGTCACGCTCGTCGTT;

Gly238Ser alone:

(SEQ ID NO:21) TTGCTGATAAATCTGGAGCCAGTGAGCGTGGGTCTC GCGGTA and
30 (SEQ ID NO:22) TGGCTCCAGATTATCAGCAA;

Gly238Ser and Arg241His (combined):

(SEQ ID NO:23) ATGCTCACTGGCTCCAGATTATCAGCAAT and
(SEQ ID NO:24) TCTGGAGCCAGTGAGCATGGGTCTCGCGGTATCATT; G4205A:
(SEQ ID NO:25) AACCTGTCCTGGCCACCATGGCCTAAATACAATCAAA
35 TATGTATCCGCTTATGAGACAATAACCCTGATA.

These separate PCR fragments were gel purified away from the synthetic oligonucleotides. 10 ng of each fragment were combined

and a reassembly reaction was performed at 94°C for 1 minute and then 25 cycles; [94°C for 30 sec, 50°C for 30 seconds and 72°C for 45 seconds]. PCR was performed on the reassembly product for 25 cycles in the presence of the *Sfi*I-containing outside primers (primers C and D from Example 5). The DNA was digested with *Sfi*I and inserted into the wild-type pUC182Sfi vector. The following mutant combinations were obtained (Table 4).

Table 4

Name	Genotype	MIC	Source of MIC
TEM-1	Wild-type	0.02	
	Glu104Lys	0.08	10
	Gly238Ser	0.16	10
TEM-15	Glu104Lys/Gly238Ser*	10	
TEM-3	Glu104Lys/Gly238Ser/Gln39Lys	10 2-32	37, 15
ST-4	Glu104Lys/Gly238Ser/Met182Thr*	10	
	Glu104Lys/Gly238Ser/Met182Thr/Ala18Val/T3959A/G3713A/G3934A/A3689G*	320	
	Glu104Lys/Gly238Ser/Met182Thr/Ala42Gly/Gly92Ser/Arg241His/T3842C/A3767G*	640	
ST-3	Glu104Lys/Gly238Ser/Met182Thr/Ala42Gly/Gly92Ser/Arg241His*	640	

* All of these mutants additionally contain the G4205A promoter mutation.

It was concluded that conserved mutations account for 9 of 15 doublings in the MIC.

Glu104Lys alone was shown to result only in a doubling of the MIC to 0.08 µg/ml, and Gly238Ser (in several contexts with one additional amino acid change) resulted only in a MIC of 0.16 µg/ml (26). The double mutant Glu104Lys/Gly238Ser has a MIC of 10 µg/ml. This mutant corresponds to TEM-15.

These same Glu104Lys and Gly238Ser mutations, in combination with Gln39Lys (TEM-3) or Thr263Met (TEM-4) result in a high level

of resistance (2-32 µg/ml for TEM-3 and 8-32 µg/ml for TEM-4 (34, 35).

A mutant containing the three amino acid changes that were conserved after the backcross (Glu104Lys/Met182Thr/Gly238Ser) 5 also had a MIC of 10 µg/ml. This meant that the mutations that each of the new selected mutants had in addition to the three known mutations were responsible for a further 32 to 64-fold increase in the resistance of the gene to cefotaxime.

The naturally occurring, clinical TEM-1-derived enzymes 10 (TEM-1-19) each contain a different combination of only 5-7 identical mutations (reviews). Since these mutations are in well separated locations in the gene, a mutant with high cefotaxime resistance cannot be obtained by cassette mutagenesis of a single area. This may explain why the maximum MIC that was obtained by 15 the standard cassette mutagenesis approach is only 0.64 µg/ml (26). For example, both the Glu104Lys as well as the Gly238Ser mutations were found separately in this study to have MICs below 0.16 µg/ml. Use of DNA shuffling allowed combinatoriality and thus the Glu104Lys/Gly238Ser combination was found, with a MIC 20 of 10 µg/ml.

An important limitation of this example is the use of a single gene as a starting point. It is contemplated that better combinations can be found if a large number of related, naturally occurring genes are shuffled. The diversity that is present in 25 such a mixture is more meaningful than the random mutations that are generated by mutagenic shuffling. For example, it is contemplated that one could use a repertoire of related genes from a single species, such as the pre-existing diversity of the immune system, or related genes obtained from many different 30 species.

Example 7. Improvement of antibody A10B by DNA shuffling of a library of all six mutant CDRs.

The A10B scFv antibody, a mouse anti-rabbit IgG, was a gift 35 from Pharmacia (Milwaukee WI). The commercially available Pharmacia phage display system was used, which uses the pCANTAB5 phage display vector.

The original A10B antibody reproducibly had only a low avidity, since clones that only bound weakly to immobilized antigen (rabbit IgG), (as measured by phage ELISA (Pharmacia assay kit) or by phage titer) were obtained. The concentration 5 of rabbit IgG which yielded 50% inhibition of the A10B antibody binding in a competition assay was 13 picomolar. The observed low avidity may also be due to instability of the A10B clone.

The A10B scFv DNA was sequenced (United States Biochemical Co., Cleveland OH) according to the manufacturer's instructions. 10 The sequence was similar to existing antibodies, based on comparison to Kabat (33).

1) Preparation of phage DNA

Phage DNA having the A10B wild-type antibody gene (10 μ l) was incubated at 99°C for 10 min, then at 72°C for 2 min. PCR 15 mix (50 mM KCl, 10 mM Tris-HCl pH 9.0, 0.1% Triton X-100, 200 μ M each dNTP, 1.9 mM MgCl₂), 0.6 μ m of each primer and 0.5 μ l Taq DNA Polymerase (Promega, Madison WI) was added to the phage DNA. A PCR program was run for 35 cycles of [30 seconds at 94°C, 30 seconds at 45°C, 45 seconds at 72°C]. The primers used were: 20 5' ATGATTACGCCAAGCTTT 3' (SEQ ID NO:26) and 5' TTGTCGTCTTCCAGACGTT 3' (SEQ ID NO:27).

The 850 bp PCR product was then electrophoresed and purified from a 2% low melting point agarose gel.

2) Fragmentation

25 300 ng of the gel purified 850 bp band was digested with 0.18 units of DNase I (Sigma, St. Louis MO) in 50 mM Tris-HCl pH 7.5, 10 mM MgCl₂ for 20 minutes at room temperature. The digested DNA was separated on a 2% low melting point agarose gel and bands between 50 and 200 bp were purified from the gel.

30 3) Construction of Test Library

The purpose of this experiment was to test whether the insertion of the CDRs would be efficient.

The following CDR sequences having internal restriction enzyme sites were synthesized. "CDR H" means a CDR in the heavy 35 chain and "CDR L" means a CDR in the light chain of the antibody.

CDR oligos with restriction sites:**CDR H1 (SEQ ID NO:34)**

5' TTCTGGCTACATCTTCACAGAATTCACTAGATTGGGTGAGGCAGACGCCTGAA3'

CDR H2 (SEQ ID NO:35)5 5' ACAGGGACTTGAGTGGATTGGAATCACAGTCAGCTTATCCTTATCTCAGGTCTCGAGT
TCCAAGTACTAAAGGCCACACTGAGTGT A 3'**CDR H3 (SEQ ID NO:36)**5' TGTCTATTCCTGTGCTAGATCTGACTGCAGTCTTATACGAGGATCCATTGGGCCAAGG
GACCAGGTCA 3'**10 CDR L1 (SEQ ID NO:37)**5' AGAGGGTCACCATGACCTGCGGACGTCTTAAGCGATCGGCTGATGGCCTGGTACCAAC
AGAACGCTGGAT 3'**CDR L2 (SEQ ID NO:38)**5' TCCCCCAGACTCCTGATTTATTAAGGGAGATCTAACAGCTGTTGGTCCCTTCGCTTCAGT
15 3'**CDR L3 (SEQ ID NO:39)**5' ATGCTGCCACTTATTACTGCTTCTGCGCGCTTAAAGGATATCTTCATTCGGAGGGGGGA
CCAAGCT 3'

The CDR oligos were added to the purified A10B antibody DNA fragments of between 50 to 200 bp from step (2) above at a 10 fold molar excess. The PCR mix (50 mM KCl, 10 mM Tris-HCl pH 9.0, 0.1% Triton x-100, 1.9 mM MgCl₂, 200 μM each dNTP, 0.3 μl Taq DNA polymerase (Promega, Madison WI), 50 μl total volume) was added and the shuffling program run for 1 min at 94°C, 1 min at 25 72°C, and then 35 cycles: 30 seconds at 94°C, 30 seconds at 55°C, 30 seconds at 72°C.

1 μl of the shuffled mixture was added to 100 μl of a PCR mix (50 mM KCl, 10 mM Tris-HCl pH 9.0, 0.1% Triton X-100, 200 μM each dNTP, 1.9 mM MgCl₂, 0.6 μM each of the two outside primers 30 (SEQ ID NO:26 and 27, see below), 0.5 μl Taq DNA polymerase) and the PCR program was run for 30 cycles of [30 seconds at 94°C, 30 seconds at 45°C, 45 seconds at 72°C]. The resulting mixture of DNA fragments of 850 basepair size was phenol/chloroform extracted and ethanol precipitated.

35 The outside primers were:

Outside Primer 1: SEQ ID NO:27

5' TTGTCGTCTTCAGACGTT 3'

Outside Primer 2: SEQ ID NO:26

5' ATGATTACGCCAAGCTTT 3'

The 850 bp PCR product was digested with the restriction enzymes *Sfi*I and *Not*I, purified from a low melting point agarose 5 gel, and ligated into the pCANTAB5 expression vector obtained from Pharmacia, Milwaukee WI. The ligated vector was electroporated according to the method set forth by Invitrogen (San Diego CA) into TG1 cells (Pharmacia, Milwaukee WI) and plated for single colonies.

10 The DNA from the resulting colonies was added to 100 μ l of a PCR mix (50 mM KCl, 10 mM Tris-HCl pH 9.0, 0.1% Triton X-100, 200 μ M each dNTP, 1.9 mM MgCl₂, 0.6 μ M of Outside primer 1 (SEQ ID No. 27; see below) six inside primers (SEQ ID NOS:40-45; see below), and 0.5 μ l Taq DNA polymerase) and a PCR program was run 15 for 35 cycles of [30 seconds at 94°C, 30 seconds at 45°C, 45 seconds at 72°C]. The sizes of the PCR products were determined by agarose gel electrophoresis, and were used to determine which CDRs with restriction sites were inserted.

CDR Inside Primers:

20 H 1 (SEQ ID NO:40) 5' AGAATTCATCTAGATTTG 3',
H 2 (SEQ ID NO:41) 5' GCTTATCCTTATCTCAGGTC 3',
H 3 (SEQ ID NO:42) 5' ACTGCAGTCTTATACGAGGAT 3'
L 1 (SEQ ID NO:43) 5' GACGTCTTAAGCGATCG 3',
L 2 (SEQ ID NO:44) 5' TAAGGGAGATCTAACACAG 3',
25 L 3 (SEQ ID NO:45) 5' TCTGCGCGCTTAAAGGAT 3'

The six synthetic CDRs were inserted at the expected locations in the wild-type A10B antibody DNA (Figure 7). These studies showed that, while each of the six CDRs in a specific clone has a small chance of being a CDR with a restriction site, 30 most of the clones carried at least one CDR with a restriction site, and that any possible combination of CDRs with restriction sites was generated.

4) Construction of Mutant Complementarity Determining Regions ("CDRs")

35 Based on our sequence data six oligonucleotides corresponding to the six CDRs were made. The CDRs (Kabat definition) were synthetically mutagenized at a ratio of 70

(existing base):10:10:10, and were flanked on the 5' and 3' sides by about 20 bases of flanking sequence, which provide the homology for the incorporation of the CDRs when mixed into a mixture of unmutagenized antibody gene fragments in a molar excess. The resulting mutant sequences are given below.

5 **oligos for CDR Library**

CDR H1 (SEQ ID NO:28)

5' TTCTGGCTACATCTTCACAACTTATGATATAGACTGGGTGAGGCAGACGCCCTGAA 3'

CDR H2 (SEQ ID NO:29)

10 5' ACAGGGACTTGAGTGGATTGGATGGATTTTCCTGGAGAGGGTGGTACTGAATACAATGA
GAAGTTCAAGGGCAGGCCACACTGAGTGTA 3'

CDR H3 (SEQ ID NO:30)

5' TGTCTATTCTGTGCTAGAGGGGACTACTATAGGCGCTACTTGACTTGAGGGCCAAGG
GACCACGGTCA 3'

15 CDR L1 (SEQ ID NO:31)

5' AGAGGGTCACCATGACCTGCAGTGCCAGCTCAGGTATACGTTACATATATTGGTACCAAC
AGAACCTGGAT 3'

CDR L2 (SEQ ID NO:32)

5' TCCCCCAGACTCCTGATTTATGACACATCCAACGTGGCTCCTGGAGTCCCTTTCGCTTCAGT
20 3'

C D R L 3 (S E Q I D N O : 3 3)
5' ATGCTGCCACTTATTACTTGCCAGGAGTGGAGTGGTTATCCGTACACGTTCGGAGGGGGGG
ACCAAGCT 3' .

Bold and underlined sequences were the mutant sequences
25 synthesized using a mixture of nucleosides of 70:10:10:10 where
70% was the wild-type nucleoside.

A 10 fold molar excess of the CDR mutant oligos were added
to the purified A10B antibody DNA fragments between 50 to 200 bp
in length from step (2) above. The PCR mix (50 mM KCl, 10 mM
30 Tris-HCl pH 9.0, 0.1% Triton x-100, 1.9 mM MgCl₂, 200 μM each
dNTP, 0.3 μl Taq DNA polymerase (Promega, Madison WI), 50 μl
total volume) was added and the shuffling program run for 1 min
at 94°C, 1 min at 72°C, and then 35 cycles: [30 seconds at 94°C,
30 seconds at 55°C, 30 seconds at 72°C].

35 1 μl of the shuffled mixture was added to 100 μl of a PCR
mix (50 mM KCl, 10 mM Tris-HCl pH 9.0, 0.1% Triton X-100, 200 μM
each dNTP, 1.9 mM MgCl₂, 0.6 μM each of the two outside primers

(SEQ ID NO:26 and 27, see below), 0.5 μ l Taq DNA polymerase) and the PCR program was run for 30 cycles of [30 seconds at 94°C, 30 seconds at 45°C, 45 seconds at 72°C]. The resulting mixture of DNA fragments of 850 basepair size was phenol/chloroform extracted and ethanol precipitated.

5 The outside primers were:

Outside Primer 1: SEQ ID NO:27 5' TTGTCGTCTTCCAGACGTT 3'

Outside Primer 2: SEQ ID NO:26 5' ATGATTACGCCAAGCTTT 3'

5) Cloning of the scFv antibody DNA into pCANTAB5

10 The 850 bp PCR product was digested with the restriction enzymes *Sfi*I and *Not*I, purified from a low melting point agarose gel, and ligated into the pCANTAB5 expression vector obtained from Pharmacia, Milwaukee WI. The ligated vector was electroporated according to the method set forth by Invitrogen 15 (San Diego CA) into TG1 cells (Pharmacia, Milwaukee WI) and the phage library was grown up using helper phage following the guidelines recommended by the manufacturer.

The library that was generated in this fashion was screened for the presence of improved antibodies, using six cycles of 20 selection.

6) Selection of high affinity clones

15 wells of a 96 well microtiter plate were coated with Rabbit IgG (Jackson Immunoresearch, Bar Harbor ME) at 10 μ g /well for 1 hour at 37°C, and then blocked with 2% non-fat dry milk in 25 PBS for 1 hour at 37°C.

100 μ l of the phage library (1×10^{10} cfu) was blocked with 100 μ l of 2% milk for 30 minutes at room temperature, and then added to each of the 15 wells and incubated for 1 hour at 37°C.

Then the wells were washed three times with PBS containing 30 0.5% Tween-20 at 37°C for 10 minutes per wash. Bound phage was eluted with 100 μ l elution buffer (Glycine-HCl, pH 2.2), followed by immediate neutralization with 2M Tris pH 7.4 and transfection for phage production. This selection cycle was repeated six times.

35 After the sixth cycle, individual phage clones were picked and the relative affinities were compared by phage ELISA, and the specificity for the rabbit IgG was assayed with a kit from

Pharmacia (Milwaukee WI) according to the methods recommended by the manufacturer.

The best clone has an approximately 100-fold improved expression level compared with the wild-type A10B when tested by 5 the Western assay. The concentration of the rabbit IgG which yielded 50% inhibition in a competition assay with the best clone was 1 picomolar. The best clone was reproducibly specific for rabbit antigen. The number of copies of the antibody displayed by the phage appears to be increased.

10

Example 8. In vivo recombination via direct repeats of partial genes

A plasmid was constructed with two partial, inactive copies of the same gene (*beta-lactamase*) to demonstrate that 15 recombination between the common areas of these two direct repeats leads to full-length, active recombinant genes.

A pUC18 derivative carrying the bacterial TEM-1 betalactamase gene was used (Yanish-Perron et al., 1985, Gene 33:103-119). The TEM-1 betalactamase gene ("Bla") confers 20 resistance to bacteria against approximately 0.02 µg/ml of cefotaxime. *Sfi*1 restriction sites were added 5' of the promoter and 3' of the end of the betalactamase gene by PCR of the vector sequence with two primers:

Primer A (SEQ ID NO: 46)

25 5' TTCTATTGACGGCCTGTCAGGCCTCATATATACTTTAGATTGATT 3'

PRIMER B (SEQ ID NO: 47)

5' TTGACGCACTGCCATGGTGGCCAAAATAACAAATAGGGTTCCGCGCAC

ATTT 3'

and by PCR of the *beta-lactamase* gene sequence with two other 30 primers:

Primer C (SEQ ID NO: 48)

5' AACTGACCACGGCCTGACAGGCCGGTCTGACAGTTACCAATGCTT 3'

Primer D (SEQ ID NO: 49)

5' AACCTGTCTGCCACCATGGCCTAAATACATTCAAATATGTAT 3'

35 The two reaction products were digested with *Sfi*1, mixed, ligated and used to transform competent *E. coli* bacteria by the procedure described below. The resulting plasmid was pUC182Sfi-

Bla-Sfi. This plasmid contains an *Sfi*1 fragment carrying the Bla gene and the P-3 promoter.

The minimum inhibitory concentration of cefotaxime for *E. coli* XL1-blue (Stratagene, San Diego CA) carrying pUC182Sfi-Bla-Sfi was 0.02 µg/ml after 24 hours at 37°C.

The tetracycline gene of pBR322 was cloned into pUC18Sfi-Bla-Sfi using the homologous areas, resulting in pBR322TetSfi-Bla-Sfi. The TEM-1 gene was then deleted by restriction digestion of the pBR322TetSfi-Bla-Sfi with *Ssp*I and *Fsp*I and blunt-end ligation, resulting in pUC322TetSfi-Sfi.

Overlapping regions of the TEM-1 gene were amplified using standard PCR techniques and the following primers:

Primer 2650 (SEQ ID NO: 50) 5' TTCTTAGACGTCAGGTGGCACTT 3'

Primer 2493 (SEQ ID NO: 51) 5' TTT TAA ATC AAT CTA AAG TAT 3'

15 Primer 2651 (SEQ ID NO: 52) 5' TGCTCATCCACGAGTGTGGAGAAGTGGTCCTGCAACTTAT 3,' and

Primer 2652 (SEQ ID NO: 53)

ACCACTTCTCCACACTCGTGGATGAGCACTTTAAAGTT

The two resulting DNA fragments were digested with *Sfi*1 and 20 *Bst*X1 and ligated into the *Sfi* site of pBR322TetSfi-Sfi. The resulting plasmid was called pBR322Sfi-BL-LA-Sfi. A map of the plasmid as well as a schematic of intraplasmidic recombination and reconstitution of functional beta-lactamase is shown in Figure 9.

25 The plasmid was electroporated into either TG-1 or JC8679 *E. coli* cells. *E. coli* JC8679 is RecBC sbcA (Oliner et al., 1993, NAR 21:5192). The cells were plated on solid agar plates containing tetracycline. Those colonies which grew, were then plated on solid agar plates containing 100 µg/ml ampicillin and 30 the number of viable colonies counted. The beta-lactamase gene inserts in those transformants which exhibited ampicillin resistance were amplified by standard PCR techniques using Primer 2650 (SEQ ID NO: 50) 5' TTCTTAGACGTCAGGTGGCACTT 3' and Primer 2493 (SEQ ID NO: 51) 5' TTTTAAATCAAATCTAAAGTAT 3' and the 35 length of the insert measured. The presence of a 1 kb insert indicates that the gene was successfully recombined, as shown in Fig. 9 and Table 5.

TABLE 5

Cell	Tet Colonies	Amp colonies	Colony PCR
TG-1	131	21	3/3 at 1 kb
JC8679	123	31	4/4 at 1 kb
vector control	51	0	

5 About 17-25% of the tetracycline-resistant colonies were also ampicillin-resistant and all of the Ampicillin resistant 10 colonies had correctly recombined, as determined by colony PCR. Therefore, partial genes located on the same plasmid will successfully recombine to create a functional gene.

Example 9.In vivo recombination via direct repeats of full-length genes.

A plasmid with two full-length copies of different alleles of the beta-lactamase gene was constructed. Homologous 5 recombination of the two genes resulted in a single recombinant full-length copy of that gene.

The construction of pBR322TetSfi-Sfi and pBR322TetSfi-Bla-Sfi was described above.

The two alleles of the beta-lactamase gene were constructed 10 as follows. Two PCR reactions were conducted with pUC18Sfi-Bla-Sfi as the template. One reaction was conducted with the following primers.

Primer 2650 (SEQ ID NO: 50) 5' TTCTTAGACGTCAGGTGGCACTT 3'

Primer 2649 (SEQ ID NO: 51)

15 5' ATGGTAGTCCACGAGTGTGGTAGTGACAGGCCGGTCTGACAGTTA
CCAATGCTT 3'

The second PCR reaction was conducted with the following primers:

Primer 2648 (SEQ ID NO: 54)

5' TGTCACTACCACACTCGTGGACTACCATGGCCTAAATACATTAAA
20 TATGTAT 3'

Primer 2493 (SEQ ID NO: 51) 5' TTT TAA ATC AAT CTA AAG TAT 3'

This yielded two Bla genes, one with a 5' Sfi1 site and a 3' BstX1 site, the other with a 5' BstX1 site and a 3' Sfi1 site.

25 After digestion of these two genes with BstX1 and Sfi1, and ligation into the Sfi1-digested plasmid pBR322TetSfi-Sfi, a plasmid (pBR322-Sfi-2BLA-Sfi) with a tandem repeat of the Bla gene was obtained. (See Figure 10).

The plasmid was electroporated into *E. coli* cells. The 30 cells were plated on solid agar plates containing 15 µg/ml tetracycline. Those colonies which grew, were then plated on solid agar plates containing 100 µg/ml ampicillin and the number of viable colonies counted. The Bla inserts in those transformants which exhibited ampicillin resistance were 35 amplified by standard PCR techniques using the method and primers described in Example 8. The presence of a 1 kb insert indicated that the duplicate genes had recombined, as indicated in Table 6.

TABLE 6

Cell	Tet Colonies	Amp Colonies	Colony PCR
TG-1	28	54	7/7 at 1kb
JC8679	149	117	3/3 at 1kb
vector control	51	0	

Colony PCR confirmed that the tandem repeat was efficiently recombined to form a single recombinant gene.

10

Example 10. Multiple cycles of direct repeat recombination - Interplasmidic

In order to determine whether multiple cycles of recombination could be used to produce resistant cells more quickly, multiple cycles of the method described in Example 9 were performed.

The minus recombination control consisted of a single copy of the betalactamase gene, whereas the plus recombination experiment consisted of inserting two copies of betalactamase as a direct repeat. The tetracycline marker was used to equalize the number of colonies that were selected for cefotaxime resistance in each round, to compensate for ligation efficiencies.

In the first round, pBR322Tetsfi-Bla-Sfi was digested with EcoRI and subject to PCR with a 1:1 mix (1 ml) of normal and Cadwell PCR mix (Cadwell and Joyce (1992) PCR Methods and Applications 2: 28-33) for error prone PCR. The PCR program was 70°C for 2 minutes initially and then 30 cycles of 94°C for 30 seconds, 52°C for 30 second and 72°C for 3 minutes and 6 seconds per cycle, followed by 72°C for 10 minutes.

The primers used in the PCR reaction to create the one Bla gene control plasmid were Primer 2650 (SEQ ID NO: 50) and Primer 2719 (SEQ ID NO: 55) 5' TTAAGGGATTTGGTCATGAGATT 3'. This resulted in a mixed population of amplified DNA fragments,

designated collectively as Fragment #59. These fragments had a number of different mutations.

The primers used in two different PCR reactions to create the two Bla gene plasmids were Primer 2650 (SEQ ID NO: 50) and 5 Primer 2649 (SEQ ID NO: 51) for the first gene and Primers 2648 (SEQ ID NO: 54) and Primer 2719 (SEQ ID NO: 55) for the second gene. This resulted in a mixed population of each of the two amplified DNA fragments: Fragment #89 (amplified with primers 2648 and 2719) and Fragment #90 (amplified with primers 2650 and 10 2649). In each case a number of different mutations had been introduced the mixed population of each of the fragments.

After error prone PCR, the population of amplified DNA fragment #59 was digested with *Sfi*I, and then cloned into pBR322TetSfi-Sfi to create a mixed population of the plasmid 15 pBR322Sfi-Bla-Sfi¹.

After error prone PCR, the population of amplified DNA fragments #90 and #89 was digested with *Sfi*I and *Bst*XI at 50°C, and ligated into pBR322TetSfi-Sfi to create a mixed population of the plasmid pBR322TetSfi-2Bla-Sfi¹ (Fig. 10).

20 The plasmids pBR322Sfi-Bla-Sfi¹ and pBR322Sfi-2Bla-Sfi¹ were electroporated into *E. coli* JC8679 and placed on agar plates having differing concentrations of cefotaxime to select for resistant strains and on tetracycline plates to titre.

An equal number of colonies (based on the number of colonies 25 growing on tetracycline) were picked, grown in LB-tet and DNA extracted from the colonies. This was one round of the recombination. This DNA was digested with *Ecr*I and used for a second round of error-prone PCR as described above.

After five rounds the MIC (minimum inhibitory concentration) 30 for cefotaxime for the one fragment plasmid was 0.32 whereas the MIC for the two fragment plasmid was 1.28. The results show that after five cycles the resistance obtained with recombination was four-fold higher in the presence of in vivo recombination.

35 Example 11.

In vivo recombination via electroporation of
fragments

Competent *E. coli* cells containing pUC18Sfi-Bla-Sfi were prepared as described. Plasmid pUC18Sfi-Bla-Sfi contains the standard TEM-1 beta-lactamase gene as described, *supra*.

A TEM-1 derived cefotaxime resistance gene from pUC18Sfi-5 cef-Sfi, (clone ST2) (Stemmer WPC (1994) *Nature* 370: 389-91, incorporated herein by reference) which confers on *E. coli* carrying the plasmid an MIC of 640 µg/ml for cefotaxime, was obtained. In one experiment the complete plasmid pUC18Sfi-cef-10 Sfi DNA was electroporated into *E. coli* cells having the plasmid pUC18Sfi-Bla-Sfi.

In another experiment the DNA fragment containing the cefotaxime gene from pUC18Sfi-cef-Sfi was amplified by PCR using the primers 2650 (SEQ ID NO: 50) and 2719 (SEQ ID NO: 55). The resulting 1 kb PCR product was digested into DNA fragments of 15 <100 bp by DNase and these fragments were electroporated into the competent *E. coli* cells which already contained pUC18Sfi-Bla-Sfi.

The transformed cells from both experiments were then assayed for their resistance to cefotaxime by plating the transformed cells onto agar plates having varying concentrations 20 of cefotaxime. The results are indicated in Table 7.

TABLE 7
Colonies/ Cefotaxime Concentration

	0.16	0.32	1.28	5.0	10.0
no DNA control	14				
ST-2 mutant, whole		4000	2000	800	400
ST-2 mutant, fragments		1000	120	22	7
Wildtype, whole	27				
Wildtype, fragments	18				

From the results it appears that the whole ST-2 Cef gene was inserted into either the bacterial genome or the plasmid after electroporation. Because most insertions are homologous, it is expected that the gene was inserted into the plasmid, replacing the wildtype gene. The fragments of the Cef gene from St-2 also

inserted efficiently into the wild-type gene in the plasmid. No sharp increase in cefotaxime resistance was observed with the introduction of the wildtype gene (whole or in fragments) and no DNA. Therefore, the ST-2 fragments were shown to yield much greater cefotaxime resistance than the wild-type fragments.

It was contemplated that repeated insertions of fragments, prepared from increasing resistant gene pools would lead to increasing resistance.

Accordingly, those colonies that produced increased cefotaxime resistance with the St-2 gene fragments were isolated and the plasmid DNA extracted. This DNA was amplified using PCR by the method described above. The amplified DNA was digested with DNase into fragments (<100 bp) and 2-4 µg of the fragments were electroporated into competent *E. coli* cells already containing pUC322Sfi-Bla-Sfi as described above. The transformed cells were plated on agar containing varying concentrations of cefotaxime.

As a control, competent *E. coli* cells having the plasmid pUC18Sfi-Kan-Sfi were also used. DNA fragments from the digestion of the PCR product of pUC18Sfi-cef-Sfi were electroporated into these cells. There is no homology between the kanamycin gene and the beta-lactamase gene and thus recombination should not occur.

This experiment was repeated for 2 rounds and the results are shown in Table 8.

TABLE 8

Round	Cef conc.	KAN control	Cef resistant colonies
5	1 replate	0.16-0.64 0.32	lawn 10 small
	2 Replate	10	10 100sm @ 2.5
	3	40 1280	100 sm

10 Example 12Determination of Recombination Formats

This experiment was designed to determine which format of recombination generated the most recombinants per cycle.

In the first approach, the vector pUC18Sfi-Bla-Sfi was amplified with PCR primers to generate a large and small fragment. The large fragment had the plasmid and ends having portions of the Bla gene, and the small fragment coded for the middle of the Bla gene. A third fragment having the complete Bla gene was created using PCR by the method in Example 8. The larger plasmid fragment and the fragment containing the complete Bla gene were electroporated into *E. coli* JC8679 cells at the same time by the method described above and the transformants plated on differing concentrations of cefotaxime.

In approach 2, the vector pUC18Sfi-Bla-Sfi was amplified to produce the large plasmid fragment isolated as in approach 1 above. The two fragments each comprising a portion of the complete Bla gene, such that the two fragments together spanned the complete Bla gene were also obtained by PCR. The large plasmid fragment and the two Bla gene fragments were all electroporated into competent *E. coli* JC8679 cells and the transformants plated on varying concentrations of cefotaxime.

In the third approach, both the vector and the plasmid were electroporated into *E. coli* JC8679 cells and the transformants were plated on varying concentrations of cefotaxime.

In the fourth approach, the complete Bla gene was electroporated into *E. coli* JC8679 cells already containing the vector pUCSF1-Sfi and the transformants were plated on varying concentrations of cefotaxime. As controls, the *E. coli* JC8679 5 cells were electroporated with either the complete Bla gene or the vector alone.

The results are presented in Figure 11. The efficiency of the insertion of two fragments into the vector is 100 X lower than when one fragment having the complete Bla gene is used. 10 Approach 3 indicated that the efficiency of insertion does depend on the presence of free DNA ends since no recombinants were obtained with this approach. However, the results of approach 3 were also due to the low efficiency of electroporation of the vector. When the expression vector is already in the competent 15 cells, the efficiency of the vector electroporation is not longer a factor and efficient homologous recombination can be achieved even with uncut vector.

Example 12. Kit for cassette shuffling to optimize vector performance

In order to provide a vector capable of conferring an optimized phenotype (e.g., maximal expression of a vector-encoded sequence, such as a cloned gene), a kit is provided comprising a variety of cassettes which can be shuffled, and optimized 25 shufflants can be selected. Figure 12 shows schematically one embodiment, with each loci having a plurality of cassettes. For example, in a bacterial expression system, Figure 13 shows example cassettes that are used at the respective loci. Each cassette of a given locus (e.g., all promoters in this example) 30 are flanked by substantially identical sequences capable of overlapping the flanking sequence(s) of cassettes of an adjacent locus and preferably also capable of participating in homologous recombination or non-homologous recombination (e.g., lox/cre or flp/frt systems), so as to afford shuffling of cassettes within 35 a locus but substantially not between loci.

Cassettes are supplied in the kit as PCR fragments, which each cassette type or individual cassette species packaged in a separate tube. Vector libraries are created by combining

the contents of tubes to assemble whole plasmids or substantial portions thereof by hybridization of the overlapping flanking sequences of cassettes at each locus with cassettes at the adjacent loci. The assembled vector is ligated to a predetermined gene of interest to form a vector library wherein each library member comprises the predetermined gene of interest and a combination of cassettes determined by the association of cassettes. The vectors are transferred into a suitable host cell and the cells are cultured under conditions suitable for expression, and the desired phenotype is selected.

Example 13.

Shuffling to optimize Green Fluorescent Protein (GFP) properties

Background

Green fluorescent protein ("GFP") is a polypeptide derived from an apopeptide having 238 amino acid residues and a molecular weight of approximately 27,000. GFP contains a chromophore formed from amino acid residues 65 through 67. As its name indicates, GFP fluoresces; it does not bioluminesce like luciferase. In vivo, the chromophore of GFP is activated by energy transfer from coelenterazine complexed with the photoprotein aequorin, with GFP exhibiting green fluorescence at 510 nm. Upon irradiation with blue or UV light, GFP exhibits green fluorescence at approximately 510 nm.

The green fluorescent protein (GFP) of the jellyfish *Aequorea victoria* is a very useful reporter for gene expression and regulation (Prasher et al. (1992) Gene 111: 229; Prasher et al. (1995) Trends In Genetics 11: 320; Chalfie et al. (1994) Science 263: 802, incorporated herein by reference). WO95/21191 discloses a polynucleotide sequence encoding a 238 amino acid GFP apoprotein which contains a chromophore formed from amino acids 65 through 67. WO95/21191 disclose that a modification of the cDNA for the apopeptide of *A. victoria* GFP results in synthesis of a peptide having altered fluorescent properties. A mutant GFP (S65T) resulting in a 4-6-fold improvement in excitation amplitude has been reported (Heim et al. (1994) Proc. Natl. Acad. Sci. (U.S.A.) 91: 12501).

Overview

Green fluorescent protein (GFP) has rapidly become a widely used reporter of gene regulation. However, in many organisms, particularly eukaryotes, the whole cell fluorescence signal was found to be too low. The goal was to improve the whole cell fluorescence of GFP for use as a reporter for gene regulation for *E. coli* and mammalian cells. The improvement of GFP by rational design appeared difficult because the quantum yield of GFP is already 0.7-0.8 (Ward et al. (1982) Photochem. 10 Photobiol. 35: 803) and the expression level of GFP in a standard *E. coli* construct was already about 75% of total protein.

Improvement of GFP was performed first by synthesis of a GFP gene with improved codon usage. The GFP gene was then further improved by the disclosed method(s), consisting of 15 recursive cycles of DNA shuffling or sexual PCR of the GFP gene, combined with visual selection of the brightest clones. The whole cell fluorescence signal in *E. coli* was optimized and selected mutants were then assayed to determine performance of the best GFP mutants in eukaryotic cells.

20 A synthetic gene was synthesized having improved codon usage and having a 2.8-fold improvement of the *E. coli* whole cell fluorescence signal compared to the industry standard GFP construct (Clontech, Palo Alto, CA). An additional 16-fold improvement was obtained from three cycles of sexual PCR and 25 visual screening for the brightest *E. coli* colonies, for a 45-fold improvement over the standard construct. Expressed in Chinese Hamster Ovary (CHO) cells, this shuffled mutant showed a 42-fold improvement of signal over the synthetic construct. The expression level in *E. coli* was unaltered at about 75% of 30 total protein. The emission and excitation maxima of the GFP were also unchanged. Whereas in *E. coli* most of the wildtype GFP ends up in inclusion bodies, unable to activate its chromophore, most of the mutant protein(s) were soluble and active. The three amino acid mutations thus guide the mutant protein into the 35 native folding pathway rather than toward aggregation. The results show that DNA sequence shuffling (sexual PCR) can solve

complex practical problems and generate advantageous mutant variants rapidly and efficiently.

MATERIALS AND METHODS

5

GFP gene construction

A gene encoding the GFP protein with the published sequence (Prasher et al. (1995) op.cit, incorporated herein by reference) (238 AA, 27 kD) was constructed from oligonucleotides. In contrast to the commercially available GFP construct (Clontech, Palo Alto, CA), the sequence included the Ala residue after the fMet, as found in the original cDNA clone. Fourteen oligonucleotides ranging from 54 to 85 bases were assembled as seven pairs by PCR extension. These segments were digested with restriction enzymes and cloned separately into the vector Alpha+GFP (Whitehorn et al. (1995) Bio/Technology 13: 1215, incorporated herein by reference) and sequenced. These segments were then ligated into the eukaryotic expression vector Alpha+ to form the full-length GFP construct, Alpha+GFP (Fig. 14). The resulting GFP gene contained altered Arginine codons at amino acid positions 73 (CGT), 80 (CGG), 96 (CGC) and 122 (CGT). To reduce codon bias and facilitate expression in *E. coli*, a number of other silent mutations were engineered into the sequence to create the restriction sites used in the assembly of the gene. These were S2 (AGT to AGC; to create an NheI site), K41 (AAA to AAG; HinDIII), Y74 (TAC to TAT) and P75 (CCA to CCG; BspE1), T108 (AGA to AGG; NnuI), L141 (CTC to TTG) and E142 (GAA to GAG; XbaI), S175 (TCC to AGC; BamHI) and S202 (TCG to TCC; SalI). The 5' and 3' untranslated ends of the gene contained XbaI and EcoRI sites, respectively. The sequence of the gene was confirmed by sequencing.

Other suitable GFP vectors and sequences can be obtained from the GenBank database, such as via Internet World Wide Web, as files: CVU36202, CVU36201, XXP35SGFP, XXU19282, XXU19279, XXU19277, XXU19276, AVGFP2, AVGFP1, XXU19281, XXU19280, XXU19278, AEVGFP, and XXU17997, which are incorporated herein by reference to the same extent as if the sequence files and comments were printed and inserted herein.

The XbaI-EcoRI fragment of Alpha+GFP, containing the whole GFP gene, was subcloned into the prokaryotic expression vector pBAD18 (Guzman et al. (1995) *J. Bacteriol.* **177**: 4121), resulting in the bacterial expression vector pBAD18-GFP (Fig. 5 14). In this vector GFP gene expression is under the control of the arabinose promoter/repressor (araBAD), which is inducible with arabinose (0.2%). Because this is the only construct with the original amino acid sequence, it is referred to as wildtype 10 GFP ('wt'). A GFP-expressing bacterial vector was obtained from Clontech (Palo Alto, CA), which is referred to herein as 'Clontech' construct. GFP expression from the 'Clontech' construct requires IPTG induction.

Gene shuffling and selection

An approximately 1 kb DNA fragment containing the whole GFP gene was obtained from the PBAD-GFP vector by PCR with primers 5'-TAGCGGATCCTACCTGACGC (near NheI site) and 15 5'GAAAATCTTCTCATCCG (near EcoRI site) and purified by Wizard PCR prep (Promega, Madison, WI). This PCR product was digested into random fragments with DNase I (Sigma) and 50-300 bp 20 fragments were purified from 2% low melting point agarose gels. The purified fragments were resuspended at 10-30 ng/ul in PCR mixture (Promega, Madison, WI; 0.2 mM each dNTP/2.2 mM MgCl₂/50 MM KCl/10 mM Tris-HCl, pH 9.0/0.1% Triton-X-100) with Taq DNA 25 polymerase (Promega) and assembled (without primers) using a PCR program of 35 cycles of 94°C 30s, 45°C 30s, 72°C 30s, as described in Stemmer, WPC (1994) *Nature* **370**: 389, incorporated herein by reference. The product of this reaction was diluted 40x into new PCR mix, and the full length product was amplified 30 with the same two primers in a PCR of 25 cycles of 94°C 30s, 50°C 30s, 72°C 30s, followed by 72°C for 10 min. After digestion of the reassembled product with NheI and EcoRI, this library of point-mutated and in vitro recombined GFP genes was cloned back 35 into the PBAD vector, electroporated into *E. coli* TG1 (Pharmacia), and plated on LB plates with 100 ug/ml ampicillin and 0.2 % arabinose to induce GFP expression from the arabinose promoter.

Mutant selection

Over a standard UV light box (365 nm) the 40 brightest colonies were selected and pooled. The pool of colonies was used as the template for a PCR reaction to obtain a pool of GFP genes.

5 Cycles 2 and 3 were performed identical to cycle 1. The best mutant from cycle 3 was identified by growing colonies in microtiter plates and fluorescence spectrometry of the microtiter plates.

For characterization of mutants in *E. coli*, DNA sequencing was performed on an Applied Biosystems 391 DNA sequencer.

CHO cell expression of GFP

The wildtype and the cycle 2 and 3 mutant versions of the GFP gene were transferred into the eukaryotic expression vector Alpha+ (16) as an EcoRI-XbaI fragment. The plasmids were transfected into CHO cells by electroporation of 10^7 cells in 0.8 ml with 40 μ g of plasmid at 400V and 250 μ F. Transformants were selected using 1 mg/ml G418 for 10-12 days.

20 FACS analysis was carried out on a Becton Dickinson FACSTAR Plus using an Argon ion laser tuned to 488 nm. Fluorescence was observed with a 535/30 run bandpass filter.

RESULTSCodon usage

25 *E. coli* expressing the synthetic GFP construct ('wt') with altered codon usage yielded a nearly 3-fold greater whole cell fluorescence signal than cells expressing the 'Clontech' construct (Fig. 15A). The comparison was performed at full induction and at equal OD₆₀₀. In addition to the substitution of poor arginine codons in the 'wt' construct and the N-terminal extension present in the 'Clontech' construct, the expression vectors and GFP promoters are quite different. The cause of the improved fluorescence signal is not enhanced expression level,

30 it is improved protein performance.

35

Sexual PCR

The fluorescence signal of the synthetic 'wt' GFP construct was further improved by constructing a mutant library by sexual PCR methods as described herein and in Stemmer WPC (1994) *Proc. Natl. Acad. Sci. (U.S.A.)* 91: 10747 and Stemmer WPC 5 (1994) *Nature* 370: 389, incorporated herein by reference, followed by plating and selection of the brightest colonies. After the second cycle of sexual PCR and selection, a mutant ('cycle 2') was obtained that was about 8-fold improved over 'wt', and 23-fold over the 'Clontech' construct. After the third 10 cycle a mutant ('cycle 3') was obtained which was 16-18-fold improved over the 'wt' construct, and 45-fold over the 'Clontech' construct (Fig. 15B). The peak wavelengths of the excitation and emission spectra of the mutants were identical to that of the 'wt' construct (Fig. 15B). SDS-PAGE analysis of whole cells 15 showed that the total level of the GFP protein expressed in all three constructs was unchanged, at a surprisingly high rate of about 75% of total protein (Fig. 16, panels (a) and (b)). Fractionation of the cells by sonication and centrifugation showed that the 'wt' construct contained mostly inactive GFP in 20 the form of inclusion bodies, whereas the 'cycle 3' mutant GFP remained mostly soluble and was able to activate its chromophore. The mutant genes were sequenced and the 'cycle 1' mutant was found to contain more mutations than the 'cycle 3' mutant (Fig. 25 17). The 'cycle 3' contained 3 protein mutations and 3 silent mutations relative to the 'wt' construct. Mutations F100S, M154T, and V164A involve the replacement of hydrophobic residues with more hydrophilic residues (Kyte and Doolittle, 1982). One plausible explanation is that native GFP has a hydrophobic site on its surface by which it normally binds to Aequorin, or to 30 another protein. In the absence of this other protein, the hydrophobic site may cause aggregation and prevent autocatalytic activation of the chromophore. The three hydrophilic mutations may counteract the hydrophobic site, resulting in reduced aggregation and increased chromophore activation. Pulse chase 35 experiments with whole bacteria at 37°C showed that the $T_{1/2}$ for fluorophore formation was 95 minutes for both the 'wt' and the 'cycle 3' mutant GFP.

CHO cells

Improvements in autonomous characteristics such as self-folding can be transferable to different cellular environments. After being selected in bacteria, the 'cycle 3' mutant GFP was transferred into the eukaryotic Alpha+ vector and expressed in chinese hamster ovary cells (CHO). Whereas in E. coli the 'cycle 3' construct gave a 16-18-fold stronger signal than the 'wt' construct, fluorescence spectroscopy of CHO cells expressing the 'cycle 3' mutant showed a 42-fold greater whole cell fluorescence signal than the 'wt' construct under identical conditions (Fig. 18A). FACS sorting confirmed that the average fluorescence signal of CHO cell clones expressing 'cycle 3' was 46-fold greater than cells expressing the 'wt' construct (Fig. 18B). As for the 'wt' construct, the addition of 2 mM sodium butyrate was found to increase the fluorescence signal about 4-8 fold.

Screening versus selection

These results were obtained by visual screening of approximately 10,000 colonies, and the brightest 40 colonies were picked at each cycle. Significant improvements in protein function can be obtained with relatively low numbers of variants. In view of this surprising finding, sexual PCR can be combined with high throughput screening procedures as an improved process for the optimization of the large number of commercially important enzymes for which large scale mutant selections are not feasible or efficient.

Example 14.Shuffling to Generate Improved Peptide Display Libraries

30

Background

Once recombinants have been characterized from a phage display library, polysome display library, or the like, it is often useful to construct and screen a second generation library that displays variants of the originally displayed sequence(s). However, because the number of combinations for polypeptides longer than seven residues is so great that all permutations will not generally be present in the primary library. Furthermore,

by mutating sequences, the "sequence landscape" around the isolated sequence can be examined to find local optima.

There are several methods available to the experimenter for the purposes of mutagenesis. For example, suitable methods 5 include site-directed mutagenesis, cassette mutagenesis, and error-prone PCR.

Overview

The disclosed method for generating mutations *in vitro* 10 is known as DNA shuffling. In an embodiment of DNA shuffling, genes are broken into small, random fragments with DNase I, and then reassembled in a PCR-like reaction, but typically without any primers. The process of reassembling can be mutagenic in the absence of a proof-reading polymerase, generating up to about 15 0.7% error rate. These mutations consist of both transitions and transversion, often randomly distributed over the length of the reassembled segment.

Once one has isolated a phage-displayed recombinant with desirable properties, it is generally appropriate to improve 20 or alter the binding properties through a round of molecular evolution via DNA shuffling. Second generation libraries of displayed peptides and antibodies were generated and isolated phage with improved (i.e., 3-1000 fold) apparent binding strength were produced. Thus, through repeated rounds of library 25 generation and selection it is possible to "hill-climb" through sequence space to optimal binding.

From second generation libraries, very often stronger binding species can be isolated. Selective enrichment of such phage can be accomplished by screening with lower target 30 concentrations immobilized on a microriter plate or in solution, combined with extensive washing or by other means known in the art. Another option is to display the mutagenized population of molecules at a lower valency on phage to select for molecules with higher affinity constants. Finally, it is possible to 35 screen second generation libraries in the presence of a low concentration of binding inhibitor (i.e., target, ligand) that blocks the efficient binding of the parental phage.

MethodsExemplary Mutagenesis Protocols

A form of recombinant DNA-based mutagenesis is known as oligonucleotide-mediated site-directed mutagenesis. An 5 oligonucleotide is designed such that can it base-pair to a target DNA, while differing in one or more bases near the center of the oligonucleotide. When this oligonucleotide is base-paired to the single-stranded template DNA, the heteroduplex is converted into double-stranded DNA in vitro; in this manner one 10 strand of the product will carry the nucleotide sequence specific by the mutagenic oligonucleotide. These DNA molecules are then propagated in vivo and the desired recombinant is ultimately identified among the population of transformants.

A protocol for single-stranded mutagenesis is described 15 below.

1. Prepare single-stranded DNA from M13 phage or phagemids. Isolate ~2 µg of DNA. The DNA can be isolated from a *dut⁻ung⁻* bacterial host (source) so that the recovered DNA contains uracil in place of many thymine residues.

20 2. Design an oligonucleotide that has at least 15 or 20 residues of complementarity to the coding regions flanking the site to be mutated. In the oligonucleotide, the region to be randomized can be represented by degenerate codons. If the non-complementary region is large (i.e., > 12 nucleotides), then the 25 flanking regions should be extended to ensure proper base pairing. The oligonucleotide should be synthesized with a 5'PO₄ group, as it improves the efficiency of the mutagenesis procedure; this group can also be added enzymatically with T4 polynucleotide kinase. (In an Eppendorf tube, incubate 100 ng of 30 oligonucleotide with 2 units of T4 polynucleotide kinase in 50 mM (pH 7.5), 10 mM MgCl₂, 5 mM DTT, and 0.1 mM ATP for 30 min.

35 3. Anneal the oligonucleotide with the single-stranded DNA in a 500 µl Eppendorf tube containing:
1 µg single-stranded DNA, 10 ng oligonucleotide, 20 mM TrisCl (pH 7.4), 2 mM MgCl₂, 50 mM NaCl.

4. Mix the solutions together and centrifuge the tube for a few seconds to recollect the liquid. Heat the tube in a flask

containing water heated to 70°C. After 5 min, transfer the flask to the lab bench and let it cool to room temperature slowly.

5. Take the tube out of the water bath and put it on ice.

Add the following reagents to the tube, for a total volume of
5 100 μ l: 20 mM Tris-HCl (pH 7.4), 2 mM DTT, 0.5 mM dATP, dCTP, dGTP
and dTTP, 0.4 mM ATP, 1 unit T7 DNA polymerase, 2 units T4 DNA ligase.

6. After 1 hr, add EDTA to 10 mM final concentration.

7. Take 20 μ l from the sample and run on an agarose gel.

10 Most of the single-stranded DNA should be converted to covalently-closed circular DNA. Electrophorese some controls in adjacent lanes (i.e., template, template reaction without oligonucleotide). Add T4 DNA ligase to close the double-stranded circular DNA.

15 8. Extract the remainder of the DNA (80 μ l) by phenol extraction and recover by ethanol precipitation.

9. Electroporate into ung⁺ bacteria.

10. Harvest the second generation phage by PEG precipitation.

20

Cassette mutagenesis

A convenient means of introducing mutations at a particular site within a coding region is by cassette mutagenesis. The "cassette" can be generated several different ways: A) by annealing two oligonucleotides together and converting them into double stranded DNA; B) by first amplifying segments of DNA with oligonucleotides that carry randomized sequences and then reamplifying the DNA to create the cassette for cloning; C) by first amplifying each half of the DNA segment with oligonucleotides that carry randomized sequences, and then heating the two pieces together to create the cassette for cloning; and D) by error-prone PCR. The cassettes formed by these four procedures are fixed in length and coding frame, but have codons which are unspecified at a low frequency. Thus, 30 cloning and expression of the cassettes will generate a plurality of peptides or proteins that have one or more mutant residues along the entire length of the cassette.

Typically, two types of mutagenesis scheme can be used. First, certain residues in a phage-displayed protein or peptide can be completely randomized. The codons at these positions can be NNN, NNK, or NNS which use 32 codons to encode all 20 residues. They can also be synthesized as preformed triplets or by mixing oligonucleotides synthesized by the split-resin method which together cover all 20 codons at each desired position. Conversely, a subset of codons can be used to favor certain amino acids and exclude others. Second, all of the codons in the cassette can have some low probability of being mutated. This is accomplished by synthesized oligonucleotides with bottles "spiked" with the other three bases or by altering the ratio of oligonucleotides mixed together by the split-resin method.

For mutagenesis of short regions, cassette mutagenesis with synthetic oligonucleotide is generally preferred. More than one cassette can be used at a time to alter several regions simultaneously. This approach is preferred when creating a library of mutant antibodies, where all six complementarity determining regions (CDR) are altered concurrently.

20

Random codons

1. Design oligonucleotides with both fixed and mutated positions. The fixed positions should correspond to the cloning sites and those coding regions presumed to be essential for binding or function.

2. During synthesis of the oligonucleotide, have the oligonucleotide synthesizer deliver equimolar amounts of each base for N, guanosine and cytosine for K, guanosine and thymidine for S.

30

"Spiked" codons

1. Design oligonucleotides with both fixed and mutated positions. The fixed positions should correspond to the cloning sites and those coding regions presumed to be essential for binding or function. The probability of finding n errors in an m long polynucleotide cassette synthesized with x fraction of the other three nucleotides at each position is represented by:

$$P = [m! / (m-n)n!] [x^n] [1-x]^{m-n}$$

2. During synthesis of the oligonucleotide switch out the base bottles. Use bottles with 100% of each base for the fixed positions and bottle with 100-x% of one base and x/3% of each of the other three bases. The doping ratio can also differ based on
5 the average amino acid use in natural globular proteins or other algorithms. There is a commercially available computer program, CyberDope, which can be used to aid in determining the base mixtures for synthesizing oligonucleotides with particular doping schemes. A demonstration copy of the CyberDope program can be
10 obtained by sending an email request to cyberdope@aol.com.

Directed codons

1. Design oligonucleotides with both fixed and mutated positions. The fixed positions should correspond to the cloning sites and those coding regions presumed to be essential for binding or function. One method has been described for inserting a set of oligonucleotides at a specific restriction enzyme site that encodes all twenty amino acids (Kegler-Ebo et al. (1994) Nucl. Acids Res. 22: 1593, incorporated herein by reference).
20 2. During synthesis of the oligonucleotide split the resin at each codon.

Error-prone PCR

There are several protocols based on altering standard PCR conditions (Saiki et al. (1988) Science 239: 487, incorporated herein by reference) to elevate the level of mutation during amplification. Addition of elevated dNTP concentrations and/or Mn⁺² increase the rate of mutation significantly. Since the mutations are theoretically introduced at random, this is one mechanism for generating populations of novel proteins. On the other hand, error-prone PCR is not well suited for altering short peptide sequences because the coding regions are short, and the rate of change would be too low to generate an adequate number of mutants for selection, nor is it ideal for long proteins,
30 35 because there will be many mutations within the coding region which complicates analysis.

1. Design oligonucleotide primers that flank the coding region of interest in the phage. They are often approximately 21 nucleotides in length and flank the region to be mutagenized. The fragment to be amplified can carry restriction sites within it to permit easy subcloning in the appropriate vector.

5 2. The following reaction is set up:

10 1 pmole of each primer; 1 pmole of the DNA template; 100 mM NaCl, 1 mM MnCl₂, 1 mM DTT, 0.2 mM of each dNTP, 2 units of Taq DNA polymerase.

15 3. Cover the liquid with mineral oil.

4. Cycle 24 times between 30 sec at 94°C, 30 sec 45°C, and 30 sec at 72°C to amplify fragments up to 1 kb. For longer fragments, the 72°C step is lengthened by approximately 30 sec for each kb.

15 5. Extend the PCR reaction for 5-10 min at 72°C to increase the fraction of molecules that are full-length. This is important if the fragment termini contain restriction sites that will be used in subcloning later.

20 6. The PCR reaction is optionally monitored by gel electrophoresis.

7. The PCR product is digested with the appropriate restriction enzyme(s) to generate sticky ends. The restriction fragments can be gel purified.

25 8. The DNA segment is cloned into a suitable vector by ligation and introduced into host cells.

DNA Shuffling

In DNA shuffling, genes are broken into small, random fragments with a phosphodiester bond lytic agent, such as DNase I, and then reassembled in a PCR-like reaction, but without requirement for any added primers. The process of reassembling can be mutagenic in the absence of a proof-reading polymerase, generating up to approximately 0.7% error when 10-50 bp fragments are used.

35 1. PCR amplify the fragment to be shuffled. Often it is convenient to PCR from a bacterial colony or plaque. Touch the colony or plaque with a sterile toothpick and swirl in a PCR

reaction mix (buffer, deoxynucleotides, oligonucleotide primers). Remove the toothpick and beat the reaction for 10 min at 99°C. Cool the reaction to 72°C, add 1-2 units of Taq DNA polymerase, and cycle the reaction 35 times for 30 sec at 94°C, 30 sec at 5 45°C, 30 sec at 72°C. and finally heat the sample for 5 min at 72°C. (Given conditions are for a 1 kb gene and are modified according the the length of the sequence as described.)

2. Remove the free primers. Complete primer removal is important.

10 3. Approximately 2-4 µg of the DNA is fragmented with 0.15 units of DNase I (Sigma, St. Louis, MO) in 100 µl of 50 mM Tris-HCl (pH 7.4), 1 mM MgCl₂, for 5-10 min at room temperature. Freeze on dry ice, check size range of fragments on 2% low melting point agarose gel or equivalent, and thaw to continue 15 digestion until desired size range is used. The desired size range depends on the application; for shuffling of a 1 kb gene, fragments of 100-300 bases are normally adequate.

4. The desired DNA fragment size range is gel purified from a 2% low melting point agarose gel or equivalent. A 20 preferred method is to insert a small piece of Whatman DE-81 ion-exchange paper just in front of the DNA, run the DNA into the paper, put the paper in 0.5 ml 1.2 M NaCl in TE, vortex 30 sec, then carefully spin out all the paper, transfer the supernatant and add 2 volumes of 100% ethanol to precipitate the DNA; no 25 cooling of the sample should be necessary. The DNA pellet is then washed with 70% ethanol to remove traces of salt.

5. The DNA pellet is resuspended in PCR mix (Promega, Madison, WI) containing 0.2 mM each DNTP, 2.2 mM MgCl₂, 50 mM KCl, 30 10 mM Tris-HCl, pH 9.0, 0.1% Triton X100, at a concentration of about 10-30 ng of fragments per µl of PCR mix (typically 100-600 ng per 10-20 µl PCR reaction). Primers are not required to be added in this PCR reaction. Taq DNA polymerase (Promega, Madison, WI) alone can be used if a substantial rate of mutagenesis (up to 0.7% with 10-50 bp DNA fragments) is desired. 35 The inclusion of a proof-reading polymerase, such as a 1:30 (vol/vol) mixture of Taq and Pfu DNA polymerase (Stratagene, San Diego, CA) is expected to yield a lower error rate and allows the

PCR of very long sequences. A program of 30-45 cycles of 30 sec 94°C, 30 sec 45-50°C, 30 sec 72°C, hold at 4°C is used in an MJ Research PTC-150 minicycler (Cambridge, MA). The progress of the assembly can be checked by gel analysis. The PCR product at this 5 point contains the correct size product in a smear of larger and smaller sizes.

6. The correctly reassembled product of this first PCR is amplified in a second PCR reaction which contains outside primers. Aliquots of 7.5 µl of the PCR reassembly are diluted 10 40x with PCR mix containing 0.8 pM of each primer. A PCR program of 20 cycles of 30 sec 94°C, 30 sec 50°C, and 30-45 sec at 72°C is run, with 5 min at 72°C at the end.

7. The desired PCR product is then digested with terminal restriction enzymes, gel purified, and cloned back into a vector, 15 which is often introduced into a host cell.

Site-specific recombination can also be used, for example, to shuffle heavy and light antibody chains inside infected bacterial cells as a means of increasing the binding affinity and specificity of antibody molecules. It is possible 20 to use the Cre/lox system (Waterhouse et al. (1993) Nucl. Acids Res. 21: 2265; Griffiths et al. (1994) EMBO J. 13: 3245, incorporated by reference) and the int system.

It is possible to take recombinants and to shuffle them together to combine advantageous mutations that occur on 25 different DNA molecules and it is also possible to take a recombinant displayed insert and to "backcross" with parental sequences by DNA shuffling to remove any mutations that do not contribute to the desired traits.

30 Example 15.

Shuffling to Generate Improved Arsenate Detoxification Bacteria

Arsenic detoxification is important for goldmining of arsenopyrite containing gold ores and other uses, such as environmental remediation. Plasmid pGJ103, containing an operon 35 encoding arsenate detoxification operon (Wang et al. (1989) Bacteriol. 171: 83, incorporated h rein by reference), was obtained from Prof. Simon Silver (U. of Illinois, Chicago, IL).

E. coli TG1 containing pJG103, containing the p1258 ars operon cloned into pUC19, had a MIC (minimum inhibitory concentration) of 4 µg/ml on LB amp plates. The whole 5.5 kb plasmid was fragmented with DNase I into fragments of 100-1000 bp, and 5 reassembled by PCR using the Perkin Elmer XL-PCR reagents. After assembling, the plasmid was digested with the unique restriction enzyme BamHI. The full length monomer was purified from the agarose gel, ligated and electroporated into E. coli TG1 cells. The transformed cells were plated on a range of sodium arsenate 10 concentrations (2, 4, 8, 16 mM in round 1), and approx. 1000 colonies from the plates with the highest arsenate levels were pooled by scraping the plates. The cells were grown in liquid in the presence of the same concentration of arsenate, and plasmid was prepared from this culture. Round 2 and 3 were 15 identical to round 1, except that the cells were plated at higher arsenate levels. 8, 16, 32, 64 mM were used for round 2; and 32, 64, 128, 256 mM were used for selection of round 3.

The best mutants grew overnight at up to 128 mM arsenate (MIC=256), a 64-fold improvement. One of the improved 20 strains showed that the TG1 (wildtype pGGJ103) grew in liquid at up to 10 mM, whereas the shuffled TG1(mutant pGJ103) grew at up to 150 mM arsenate concentration.

PCR program for the assembly was 94°C 20s, 50x(94°C 25 15s, 50°C 1 min, 72°C 30s+2s/cycle), using a circular PCR format without primers.

Four cycles of the process resulted in a 50-100-fold improvement in the resistance to arsenate conferred by the shuffled arsenate resistance operon; bacteria containing the improved operon grew on medium containing up to 500 mM arsenate.

30 Figure 19 shows enhancement of resistance to arsenate toxicity as a result of shuffling the pGJ103 plasmid containing the arsenate detoxification pathway operon.

35 Example 16. Shuffling to Generate Improved Cadmium Detoxification Bacteria

Plasmid pYW333, containing an operon for mercury detoxification is a 15.5 kb plasmid containing at least 8 genes encoding a pathway for mercury detoxification (Wang et al. (1989)

Bacteriol. 171: 83, incorporated herein by reference), was obtained from Prof. Simon Silver (Univ. Illinois, Chicago, IL). 400-1500 bp fragments were obtained as described supra and assembled with the XL-PCR reagents. After direct electroporation 5 of the assembled DNA into to *E. coli* TG1, the cells were plated on a range of levels of mercury chloride (Sigma) under a similar protocol as that described for arsenate in Example 15. The initial MIC of mercury was 50-70 μ M. Four cycles of whole plasmid shuffling were performed and increased the detoxification 10 measured as bacterial resistance to mercury from about 50-70 μ M to over 1000 μ M, a 15-20 fold improvement.

Example 17.

Enhancement of Shuffling Reactions by
Addition of Cationic Detergent

15 The rate of renaturation of complementary DNA strands becomes limiting for the shuffling long, complex sequences. This renaturation rate can be enhanced 10,000-fold by addition of simple cationic detergent (Pontius and Berg (1991) PNAS 88: 8237). The renaturation is specific and independent of up to a 20 10^6 -fold excess of heterologous DNA. In the presence of these agents the rate which the complementary DNA stands encounter each other in solution becomes limiting.

25 Addition of TMAC in an assembly reaction of a 15 kb plasmid followed by electroporation into *E. coli* resulted in the following results:

	<u>TMAC (mM)</u>	<u># Colonies</u>
	0	3
	15	88
30	30	301
	60	15
	90	3

35 Addition of CTAB in an assembly reaction of a 15 kb plasmid followed by electroporation into *E. coli* resulted in the following results:

	<u>CTAB (mM)</u>	<u># Colonies</u>
		155

0	3
30	34
100	14
300	0

5

Example 18. Sequence Shuffling via PCR Stuttering

Stuttering is fragmentation by incomplete polymerase extension of templates. A recombination format based on very short PCR extension times was employed to create partial PCR products, which continue to extend off a different template in the next (and subsequent) cycle(s). There was a strong growth rate bias between very similar templates, indicating that this format has significant limitations for the application to complex pools. If used with a rapid cycler such as the aircycler, this format may work better.

PCR programs used for PCR stuttering were 100 cycles of (94°C 15 sec, 60°C 15 sec). Two separate PCR reactions were run to obtain 1 kb PCR fragments containing the each of two GFP-negative recombination assay substrates (GFPstop1 and GFP stop2). GFP-positive recombinants can only be obtained by recombination of these two templates. The oligonucleotide primers used at the 5' end is 5'TAGCGGATCCTACCTACCTGACGC, containing an NheI site, and the oligo at the 3' end is 5'GAAAATCTTCTCTCATCC, containing an EcoRI site. A PCR reaction was set up with a 1ng of each GFP-negative gene as a template. Taq PCR reagents can be used, but the use of error-prone PCR conditions (Leung, 1989; Cadwell and Joyce, 1992) which reduce the processivity, can increase the percentage of GFP-positive recombinants.

A stuttering program of 50-150 cycles of 94°C 10-20s, 60°C 10-30s was used on a Stratagene Robocycler. The stuttered PCR product was digested with NheI and Eco RI and cloned back into pBAD18 vector digested with NheI and EcoRI and electroporated into E. coli and plated on amp plates. GFP-positive colonies arise by recombination between the two GFP-negative DNA sequences and were detected. The percentage of GFP-positive colonies obtained by stuttering was between 0.1-10%, depending on conditions.

A synthetic gene was designed for each protein, using the identical (optimal E.coli) codon usage based on the native amino acid sequence. This approach increases the DNA homology of the synthetic genes relative to the naturally occurring genes, 5 and allows us to shuffle more distantly related sequences than would be possible without the codon usage adjustment. Each of the four genes was assembled from 30 60 mer oligos and 6 40 mers. The assembly was performed by assembly PCR as described by Stemmer et al (1995; Gene 164:49-53). After assembling, the 10 genes were cloned into Sfi 1 sites of the vector pUC322 Sfi-BLA-Sfi (Stemmer et al (1995) Gene 164:49-53), and plated on a selective media. The minimum inhibitory activity of these four constructs for a wide variety of betalactam antibiotics was established. Cefotaxime was one of the antibiotics that was 15 selected for optimization against. The four genes were shuffled by pooling 1 ug of the CR product of each gene, followed by DNaseI digestion of the pool and purification of 100-300 bp fragments from agarose gels. The purified fragments were reassembled by sexual PCR initially without outside primers, and 20 then the full-length product was amplified in the presence of the outside primers. The resulting full-length genes were digested with SfiI and ligated into fresh pUC322Sfi-Sfi vector and electroporated into fresh E. coli cells, and plated on increasing concentration of several antibiotics, including cefotaxime, as 25 described previously by Stemmer (1994) Nature 370:389-391.

A manual shuffling PCR protocol is preferred for the mixing of genes that are less than 80-90% homologous. The manual PCR uses a heat-labile DNA polymerase, such as DNA polI Klenow fragment. The initial PCR program with fragments in Klenow 30 buffer at 10-30ng/ul and dNTPs:

- 1-Denature 94°C 20s
- 2-Quick-cool: dry ice ethanol 5s, ice 15s
- 3-Add Klenow enzyme
- 4-Anneal/extend 2 min 25°C
- 35 5-cycle back to denature (cycle 1)

This is repeated for 10-20 cycles to initiate the template switching, after which regular PCR with heatstable polymerases

is continued for an additional 10-20 cycles to amplify the amount of product.

Example 19.

Shuffling of Antibody Phage Display Libraries

A stable and well-expressed human single-chain Fv framework ($V_H^{251}-V_L^{A25}$) was obtained from an Ab-phage library constructed from naive human mRNA by selection for binding to diphtheria toxin. This scFv framework was used to construct a naive Ab-phage library containing six synthetically mutated CDRs based on the germline sequences. The degree of mutagenesis of each residue was similar to its naturally occurring variability within its V-region family.

A PCR product containing the scFv gene was randomly fragmented with DNaseI digestion and fragments of 50-100 bp were purified. Synthetic oligonucleotides, each containing a mutated CDR flanked by 19 bp of homology to the scFv template, were added to the random fragments at a 10:1 molar ratio. A library of full length, mutated scFv genes was reassembled from the fragments by sexual PCR. Cloning into the pIII protein of M13 phage yielded an Ab-phage library of 4×10^7 plaque-forming units. The combinations of mutant and native CDRs were characterized by colony PCR with primers specific for the native CDRs (see, Fig. 7). All six mutated CDRs were incorporated with 32-65% efficiency and a wide variety of combinations. Sequencing of the mutated CDRs showed that the observed mutation rate matched the expected rate.

This Ab-phage library was panned for two rounds in microtiter plates for binding to ten human protein targets, and seven of these targets yielded ELISA-positive clones. One target which resulted in positive clones was the human G-CSF receptor. The G-CSF receptor positive clones were subjected to a second round and one quarter of second round phage clones were ELISA-positive for binding to the G-CSF receptor.

This diverse pool was used to evaluate the suitability of three different sequence optimization strategies (conventional PCR, error-prone PCR, and DNA shuffling). After a single cycle of each of these alternatives, the DNA shuffling showed a seven-

fold advantage, both in the percentage of Ab-phage recovered and in the G-CSF receptor-specific ELISA signal. The panning was continued for six additional cycles, shuffling the pool of scFv genes after each round. The stringency of selection was 5 gradually increased to two one-hour washes at 50°C in PBS-Tween in the presence of excess soluble G-CSF receptor. In rounds 3 to 8, nearly 100 percent of the clones were ELISA positive. When pools from different cycles were assayed at identical stringency, the percentage of phage bound increased 440-fold from cycle 2 to 10 cycle 8, as shown in Fig. 31. Individual phage clones from each round showed a similar increase in specific ELISA signal. Sequencing showed that the scFv contained an average of 34 (n=4) amino acid mutations, of which only four were present in all sequences evaluated.

15 In order to reduce potential immunogenicity, neutral or weakly contributing mutations were removed by two cycles of backcrossing, with a 40-fold excess of a synthetically constructed germline scFv gene, followed by stringent panning. The average number of amino acid mutations in the backcrossed 20 scFvs were nearly halved to 18 (n=3), of which only four were present in all sequences. The backcrossed Ab phage clones were shown to bind strongly and with excellent specificity to the human G-CSF receptor. Fig. 32 shows the effect of ten selection 25 rounds for several human protein targets; six rounds of shuffling and two rounds of backcrossing were conducted. Fig. 33 shows the relative recovery rates of phage, by panning with BSA, Ab 179, or G-CSF receptor, after conventional PCR ("non-shuffled"), error-prone PCR, or recursive sequence recombination ("shuffled").

30

Example 20. Optimization of GFP in Mammalian Cells

The plasmid vector pCMV-GFP, which encodes GFP and expresses it under the control of a CMV promoter, was grown in TG1 cells and used to transfect CHO cells for transient 35 expression assays.

Plasmid was rescued from FACS selected transiently expressing TG1 cells by a proteinase K method or a PreTaq method

(Gibco/BRL). Basically, the FACS collected cells were pelleted by centrifugation, freeze-thawed repetitively, incubated with either proteinase K or PreTaq, phenol/chloroform extracted, ethanol precipitated, used to transform E. coli which were then
5 plated on Amp plates. The results were:

Proteinase K method: input - 5×10^4 rescued 3×10^4
PreTaq method: input - 5×10^4 rescued 2×10^4

The rescued plasmid was grown up and 5 µg was partially
10 digested with DNaseI and 50 to 700 bp fragments were gel purified, ethanol precipitated, and resuspended in 330µl of 3.3x PCR buffer, 44 µl Mg(OAc)₂ (Perkin-Elmer), 193 µl 40% PEG, 80 µl 10 mM dNTPs, 20 µl Tth polymerase, 2 µl Pfu polymerase, 7 µl TMAC (Sigma), and 367 µl H₂O. PCR was conducted on a MJ Research PTC-
15 150 minicycler for 40 cycles (94°C, 30 sec; 50°C, 30 sec, 72°C, 60 sec) with three sets of primers, which yielded three end-overlapping PCR fragments, which together and after digestion and ligation reconstituted the entire plasmid. The PCR fragments were digested with AlwN1 and the fragments were gel purified,
20 ligated, and electroporated into TG1 cells. Plasmid DNA was prepared and electroporated into CHO cells, which were screened by FACS for the cells transiently expressing the brightest GFP signals.

While the present invention has been described with reference to what are considered to be the preferred examples, it is to be understood that the invention is not limited to the disclosed examples. To the contrary, the invention is intended to cover various modifications and equivalent arrangements included within the spirit and scope of the appended claims.
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All publications, patents and patent applications are herein incorporated by reference in their entirety to the same extent as if each individual publication, patent or patent application

was specifically and individually indicated to be incorporated by reference in its entirety. PCT/US95/02126 filed 17 February 1995 and published on 24 August 1995 as WO95/22625 is incorporated herein by reference. Copending application attorney 5 docket no. 16528A-0196 is incorporated herein by reference.

WHAT IS CLAIMED IS:

1 1. A method of evolving a polynucleotide sequence
2 toward a desired property comprising:

3 (1) recombining at least first and second forms of the
4 polynucleotide sequence to produce a library of recombinant forms
5 of the sequence;

6 (2) screening at least a first recombinant sequence from
7 the library for evolution toward the desired property;

8 (3) recombining the first recombinant sequence with a
9 further form of the polynucleotide sequence, the same or
10 different from the first and second forms, to produce a further
11 library of recombinant polynucleotides;

12 (4) screening at least one further recombinant
13 polynucleotide from the further library of recombinant
14 polynucleotides that has further evolved toward the desired
15 property;

16 (5) repeating (3) and (4), as necessary, until the further
17 recombinant polynucleotide has acquired the desired property.

1 2. The method of claim 1, wherein at least one
2 recombining step is performed in vivo.

1 3. The method of claim 2, wherein at least two
2 recombining steps are performed in vivo.

1 4. The method of claim 2, at least one recombining
2 step is between different forms of the polynucleotide sequence
3 in separate plasmid vectors.

1 5. The method of claim 2, wherein at least one
2 recombining step is between a first form of the polynucleotide
3 sequence in a viral vector and a second form of the
4 polynucleotide sequence in a plasmid vector.

1 6. The method of claim 2, wherein at least one
2 recombining step is between different forms of the polynucleotide
3 sequences in separate viral vectors.

1 7. The method of claim 2, wherein at least one
2 recombining step is between a first form of the polynucleotide
3 and a second form of the polynucleotide in a host chromosome.

1 8. The method of claim 7, wherein the first form of
2 the polynucleotide is in a plasmid vector.

1 9. The method of claim 7, wherein the first form of
2 the polynucleotide is in a viral vector.

1 10. The method of claim 2, wherein different forms of
2 the polynucleotide are contained in a population of cells and the
3 cells are exposed to an electric field promoting exchange of the
4 different forms between the cells.

1 11. The method of claim 2, wherein different forms of
2 the polynucleotide are contained in a collection of cells and the
3 different forms are exchanged between cells by conjugation.

1 12. The method of claim 1, wherein at least one
2 recombining step is effected by nonhomologous recombination of
3 different forms of the polynucleotide.

1 13. The method of claim 1, wherein the polynucleotide
2 has introns and exons and at least one recombining step is
3 effected by homologous recombination between introns of different
4 forms of the polynucleotide.

1 14. The method of claim 1, wherein at least one
2 recombining step is performed in vivo and at least one
3 recombining step is performed in vitro.

1 15. The method of claim 1, wherein at least two
2 recombining steps are performed in vitro.

1 16. The method of claim 1, wherein at least one
2 recombining step is performed by incubating different forms of
3 the polynucleotide under conditions suitable for polymerase chain
4 replication in the presence of an agent that promotes switching
5 of the polymerase between templates, whereby the replication
6 generates recombinant forms of the different polynucleotides.

1 17. A method for forming a mutagenized double-stranded
2 polynucleotide from a template double-stranded polynucleotide,
3 wherein the template double-stranded polynucleotide has been
4 converted into double-stranded fragments of a desired size,
5 comprising:

6 a) adding to the resultant population of double-
7 stranded random fragments one or more single or double-stranded
8 oligonucleotides, wherein said oligonucleotides comprise an area
9 of structural or sequence similarity and an area of heterology
10 to the double-stranded template polynucleotide;

11 b) denaturing the resultant mixture of double-
12 stranded random fragments and oligonucleotides into single-
13 stranded fragments;

14 c) incubating the resultant population of
15 single-stranded fragments with a polymerase under conditions
16 which result in the annealing of said single-stranded fragments
17 at said areas of similarity to form pairs of annealed fragments,
18 said areas of similarity being sufficient for one member of a
19 pair to prime replication of the other thereby forming a
20 mutagenized double-stranded polynucleotide; and

21 d) repeating steps (b) and (c) for at least two
22 further cycles, wherein the resultant mixture in step (b) of a
23 further cycle includes the mutagenized double-stranded
24 polynucleotide from step (c) of the previous cycle, and the
25 further cycle forms a further mutagenized double-stranded
26 polynucleotide.

1 18. The method of claim 17, wherein the incubating is
2 performed in the presence of an agent that promotes annealing of
3 the single-stranded fragments.

1 19. The method of claim 18, wherein the agent is
2 selected from the group consisting of PEG, a cationic detergent,
3 an exonuclease or a recombinogenic protein.

1 20. The method of claim 19, wherein the agent is recA.

1 21. The method of claim 17, wherein the polynucleotide
2 is converted into double-stranded fragments of a desired size by
3 PCR stuttering.

1 22. The method of claim 17, wherein the polynucleotide
2 is converted into double-stranded fragments of a desired size by
3 replication of the polynucleotide in the presence of UTP and
4 cleavage of the replicated polynucleotide with UDG glycosylase.

1 23. The method of claim 1, further comprising
2 selecting against unrecombined substrates for recombination after
3 at least one recombining step.

1 24. The method of claim 2, wherein at least one
2 recombining step is performed in a mutator host cell or a host
3 cell exposed to a mutagen.

1 25. A method for generating an enhanced GFP protein
2 and polynucleotides encoding same, comprising performing DNA
3 shuffling on a GFP encoding expression vector and selecting or
4 screening for variants having an enhanced desired property.

1 26. The method of claim 25, wherein the method
2 comprises a step of error-prone or mutagenic amplification or
3 site-directed mutagenesis.

1 27. The method of claim 26, wherein the error-prone
2 or mutagenic amplification or site-directed mutagenesis
3 introduces a mutation in the region outside the chromophore
4 segment comprising amino acids 64-69.

1 28. The method of claim 25, wherein the property is
2 fluorescence intensity.

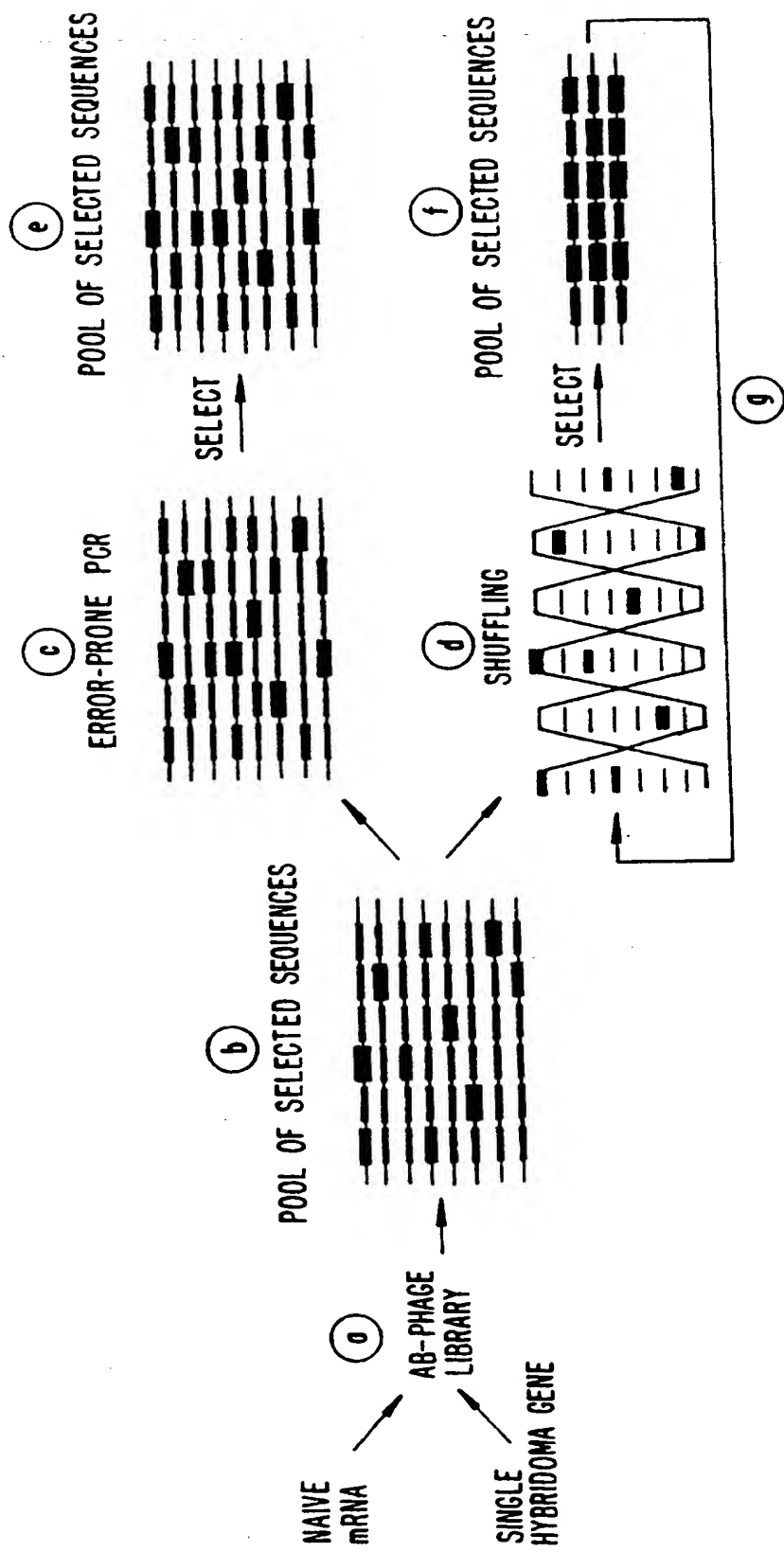
1 29. An enhanced GFP protein comprising a point
2 mutation as compared to wildtype sequence outside the chromophore
3 region comprising amino acids 64-69.

1 30. The enhanced GFP protein of claim 29, wherein the
2 mutation is in the region from amino acid 100 to amino acid 173.

1 31. The enhanced GFP protein of claim 30, wherein the
2 mutation is at residue 100, 154, and 173.

1 32. The enhanced GFP protein of claim 31, wherein the
2 mutation comprises a substitution selected from the group: F100S,
3 M154T or E173G.

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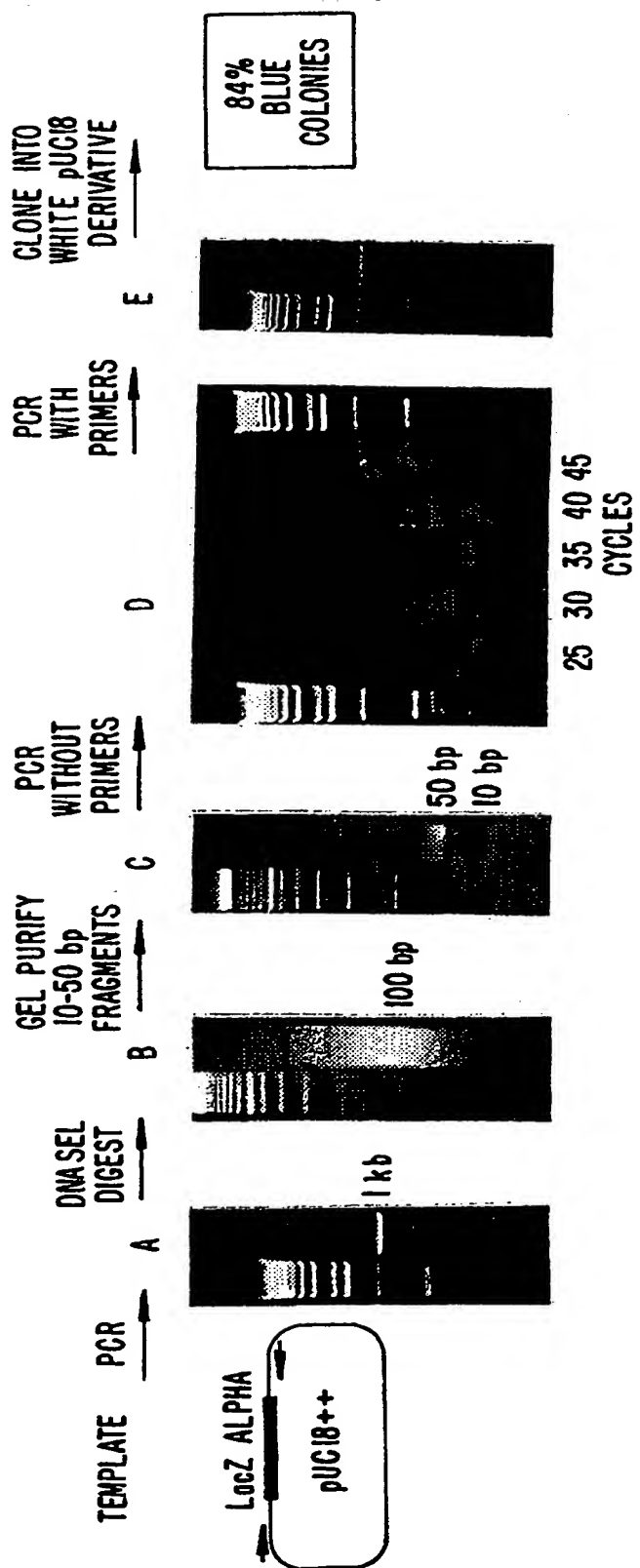
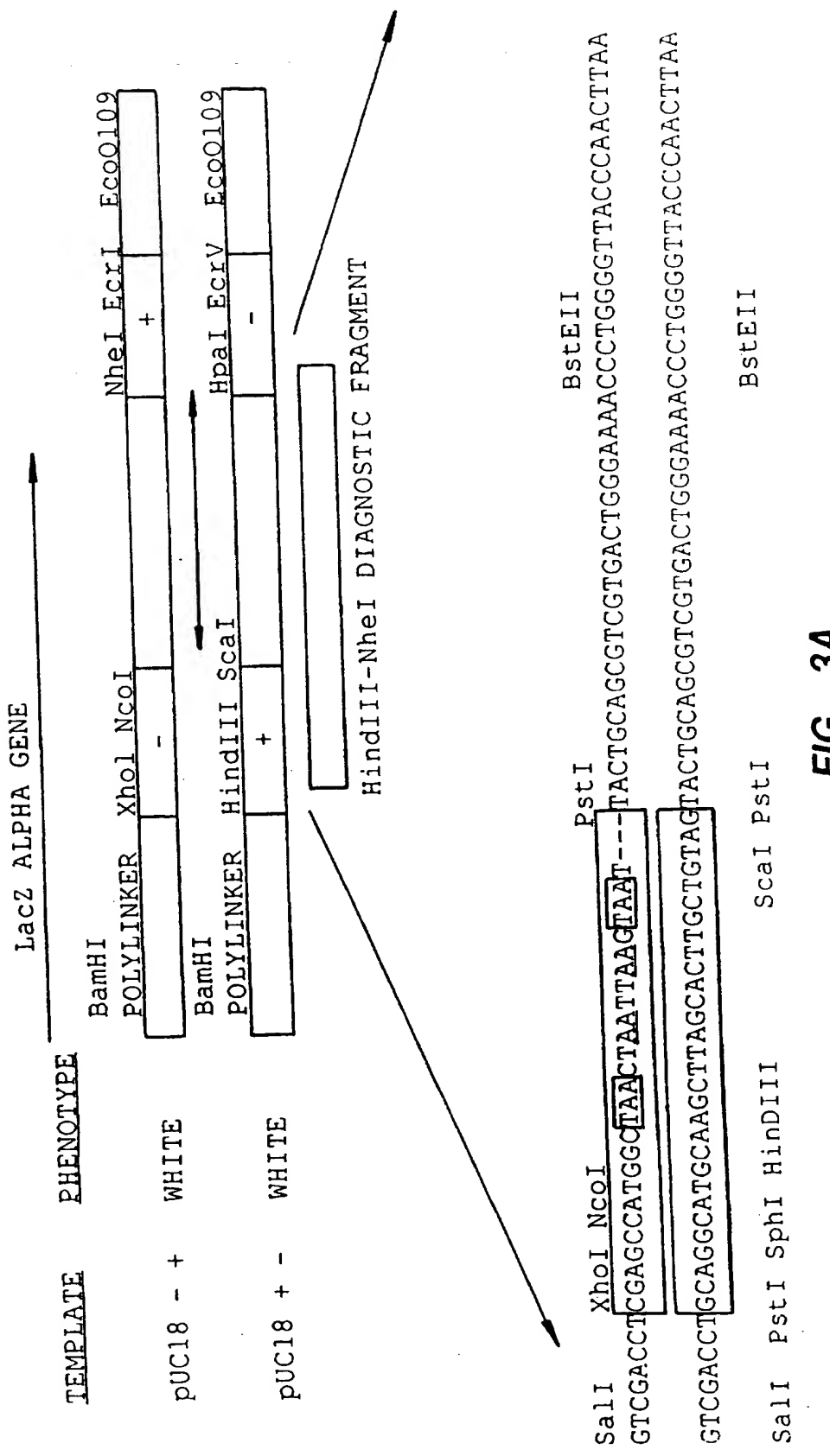


FIG. 2.

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**SUBSTITUTE SHEET (RULE 26)**

The diagram shows a vertical DNA sequence with several restriction enzyme recognition sites highlighted by boxes:

- NheI**: A box highlights the sequence **GCCTTGCATCCACCTTCGCTAGCTGGGAATTCCGAAGAA**.
- ECORI**: A box highlights the sequence **GCCTTGCATCCACCTTCGCTAGCTGGGAATTCCGAAGAA**.
- BssHII**: A box highlights the sequence **GCCTTGCATCCACCTTCGCTAGCTGGGAATTCCGAAGAA**.
- HpaI**: A box highlights the sequence **GCCTTGCATCCACCTTCGCTAGCTAACTTAAGATAATCGCGC**.
- EcrV**: A box highlights the sequence **GCCTTGCATCCACCTTCGCTAGCTAACTTAAGATAATCGCGC**.
- PspI**: A box highlights the sequence **GCCTTGCATCCACCTTCGCTAGCTAACTTAAGATAATCGCGC**.
- PvuI**: A box highlights the sequence **GCCTTGCATCCACCTTCGCTAGCTAACTTAAGATAATCGCGC**.

A small arrow points downwards from the bottom of the sequence.

FIG. 3B.

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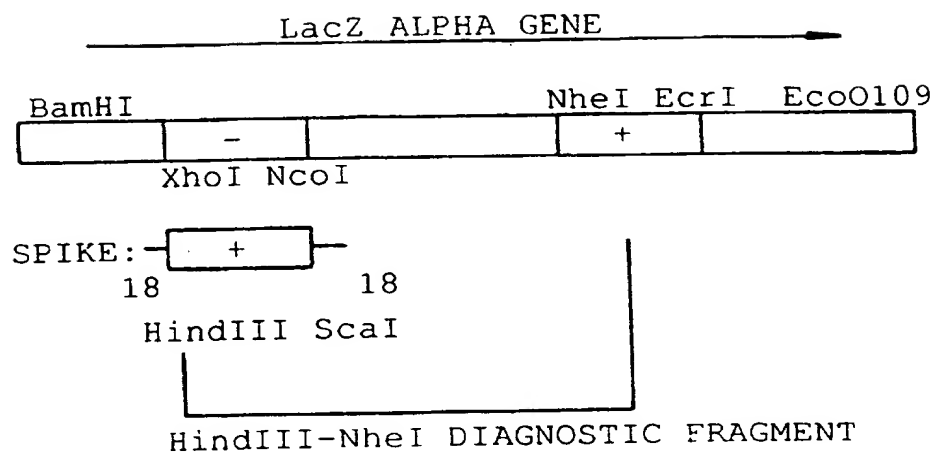


FIG. 4.

M ATGGTTCCGATCCGTAGCTGCACCTACCGTCTGGCAGAACAGCAGAAAAGCCTGGTCTGTCCGACCCGTACGAACTGAAGCT
II ATGGCACCCGTTAGATCTCTGAACTGCAACCTTCTGCGACTCCCAACAGAAAAGCTGGTCTGTCCGACCCGTACGAACTGAAGCT

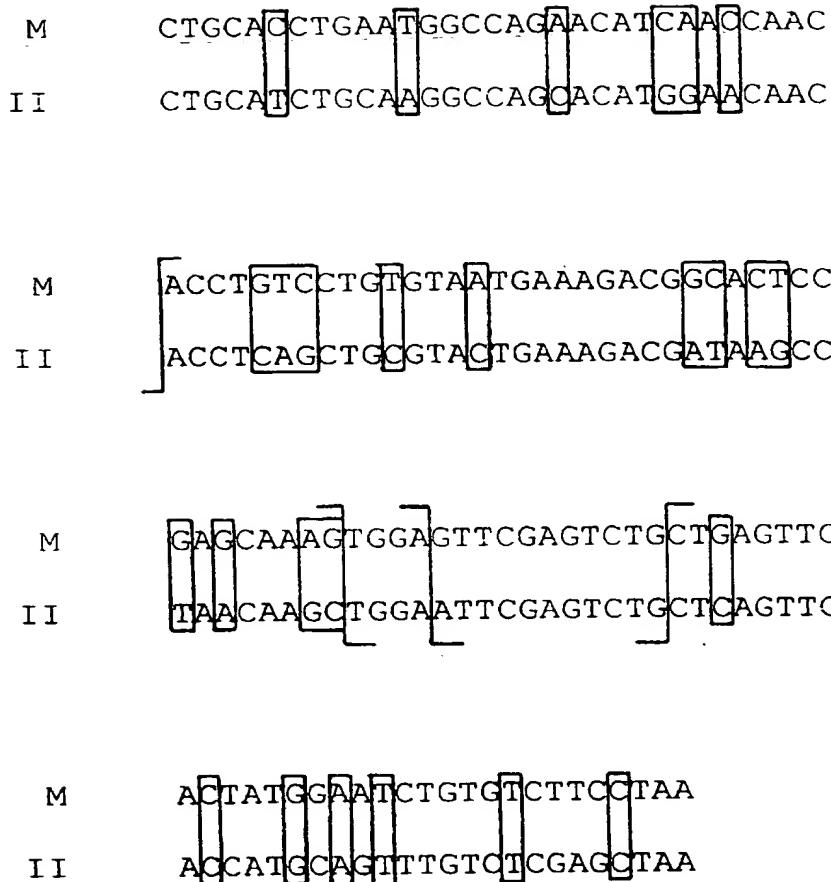
M	AGGTGAAATTCTCATGACCTTCGCTCAAGGTGAAACCAAGCAAACGACAAATCCC
I	AGGTGAAATTCTCATGACCTTCGCTCAAGGTGAAACGACAAAGATCCCAACTG

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M GAOCTCTGAGCGTGGACCCAAACAGTAACTCCAAAGAAGAATGGAGAACAGATTTCAAGATGGAAGCTCAA
II GAOCTCTGAGCGTGAATCCGTGACCCGAAAGACTATCCGAAAGAAGAAATGGAGAACAGATTTAACAAAGATTTGAGATTAA

M CCGAACTGGTACATCAGGCACTCAAGAGCCCAAAGGCCTGTTTCCCTGG . . TACAACTCCGGTCAGGATTATCGACTTCTC
II CCAAACTGGTACATCAGGCACTCAAGAGCCCAAAGGCCTGTTTCCCTGGGGTACCAAAGGGGTCAGGATTATCGACTTCTC

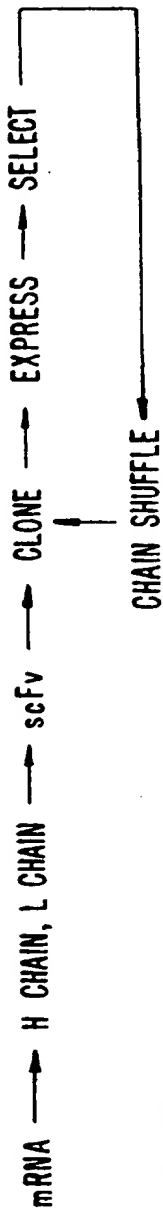
FIG. 5A.

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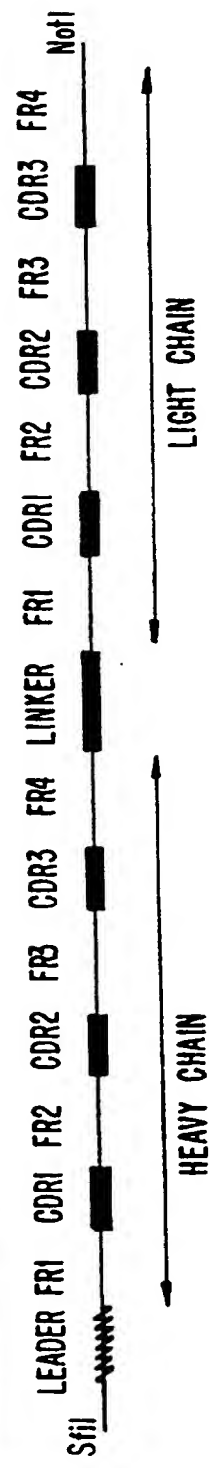
**FIG. 5B.**

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A10B = scFv OF ANTI-R-IgG ANTIBODY (PHARMACIA)



scFv STRUCTURE



TOTAL: 750 bp

FIRST EXPERIMENT:



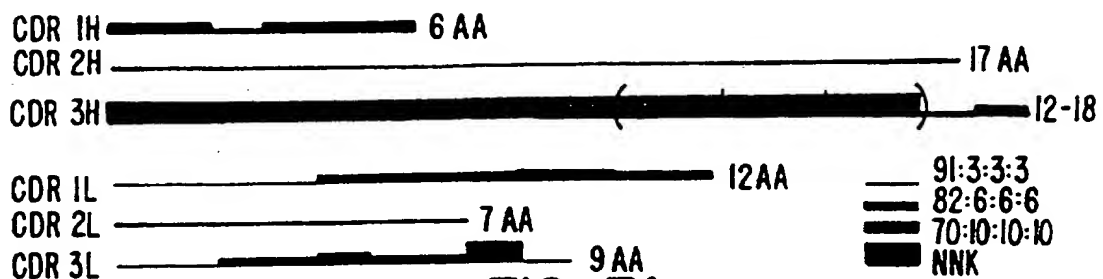
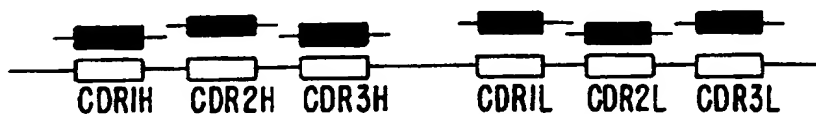
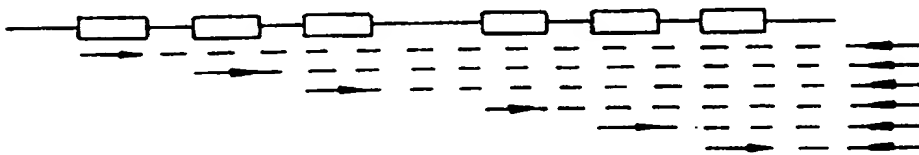
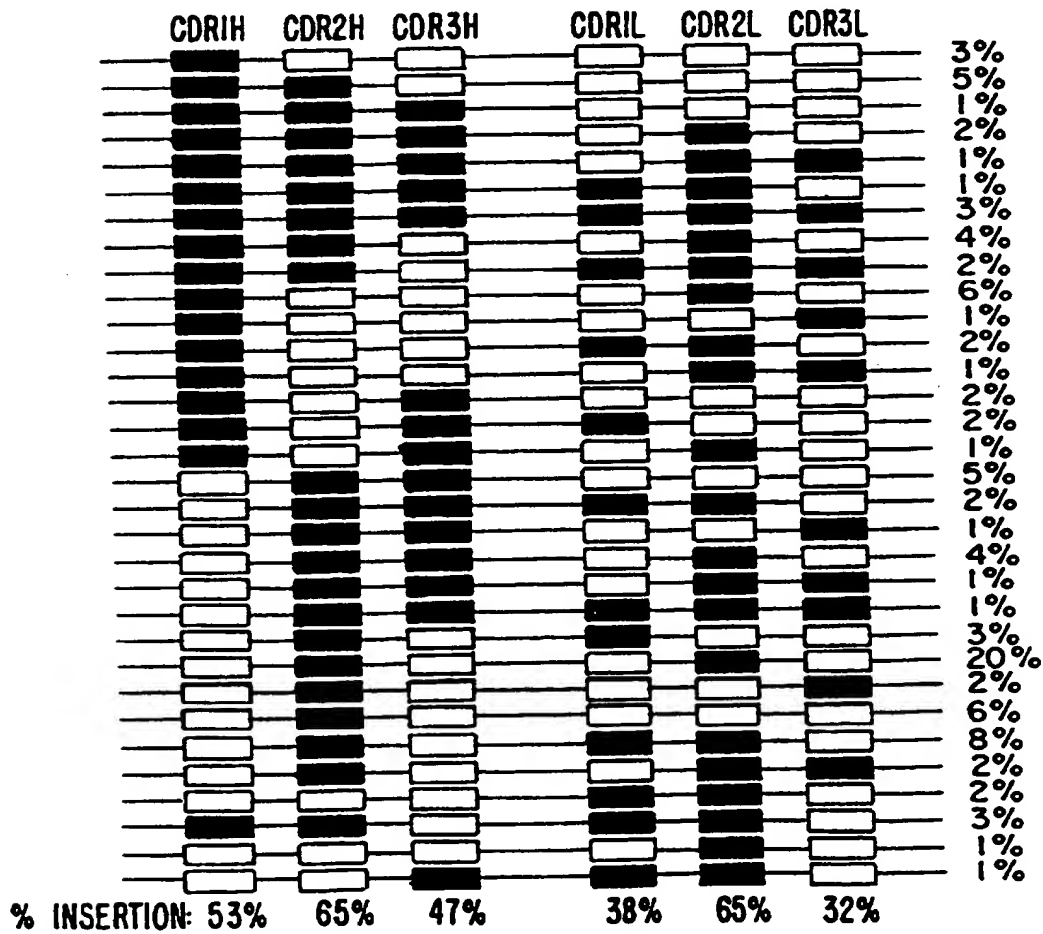
A10BscFv

SPIKE: 70/10/10/10CDRs:

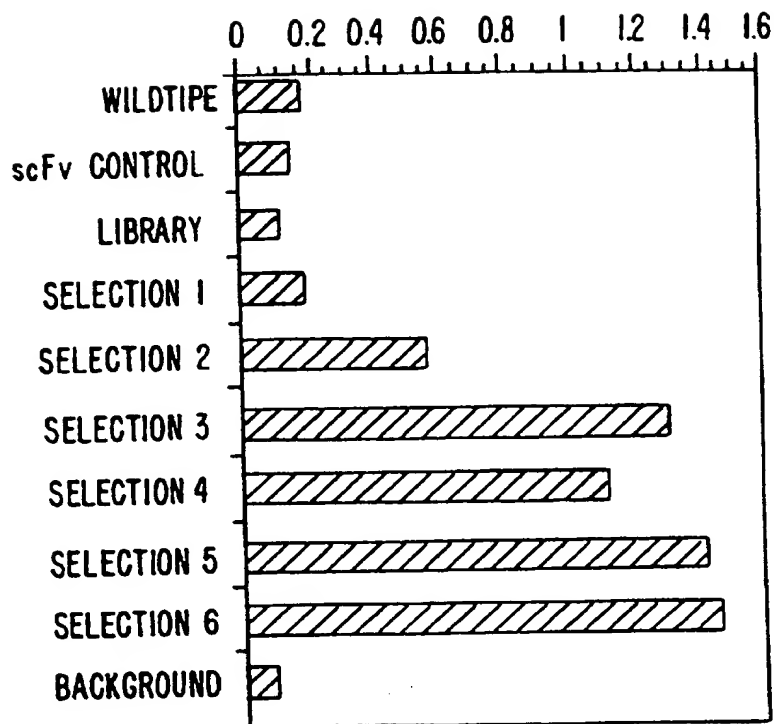


FIG. 6.

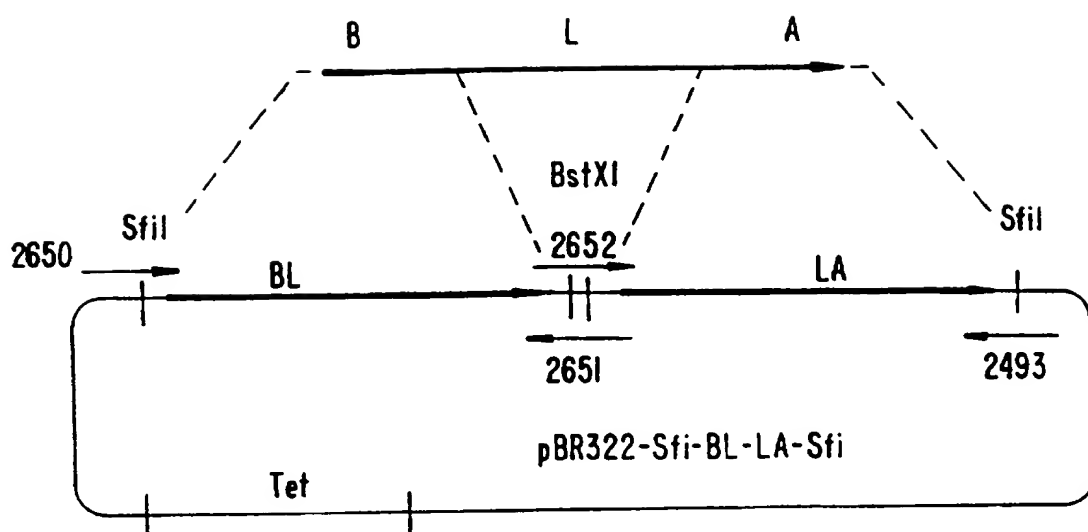
9/37

**FIG. 7A.****FIG. 7B.****FIG. 7C.****FIG. 7D.**

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 RABBIT**FIG. 8.**

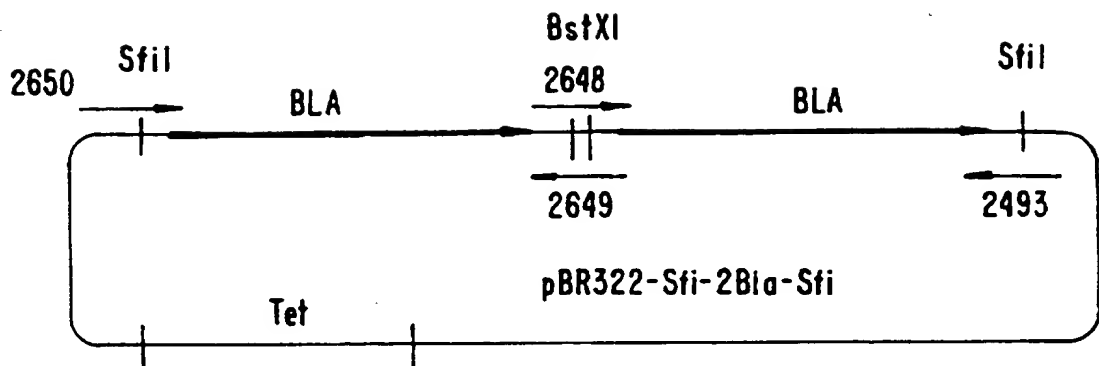
II/37



CELL	Tet COLONIES	Amp COLONIES	COLONY PCR
TG-1	131	21	3/3 AT 1 KB
JC8679	123	31	4/4 AT 1 KB
VECTOR CONTROL	51	0	

FIG. 9.

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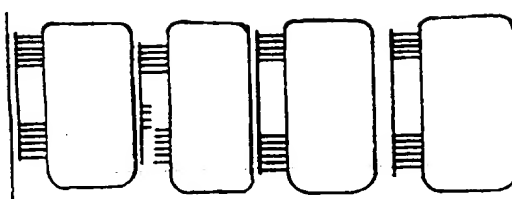
CELL	Tet COLONIES	Amp COLONIES	COLONY PCR
TG-1	28	54	7/7 AT 1 KB
JC8679	149	117	3/3 AT 1 KB
VECTOR CONTROL	51	0	

FIG. 10.

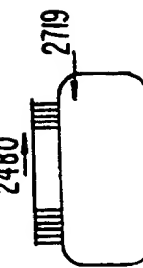
APPROACH	AMP COLONIES	TET COLONIES	% HOMOLOGOUS RECOMBINATION	COMMENT
1-CUT VECTOR	4,000	1,500	100% (N=14)	EFFICIENT INSERTION BY HOMOLOGOUS RECOMBINATION WITH CO-ELECTROPORATED VECTOR
1 INSERT				
JC8679	2,000	16	100% (N=2)	100x LESS EFFICIENT THAN 1 FRAGMENT
2-CUT VECTOR				
2 INSERTS				
JC8679	3-UNCUT VECTOR	16	0	HOMOLOGOUS INSERTION DEPENDS ON FREE ENDS
	1 INSERT			
JC8679	4-NO VECTOR	5,000	10,000	70% (N=7)
	1 INSERT			IF VECTOR IS IN CELLS ALREADY, HIGH EFFICIENCY OCCURS EVEN THROUGH VECTOR IS UNCUT
JC8679::PUCSfi-Sfi	5-NO VECTOR	2,000	0	-CONTROL: NON-HOMOLOGOUS INSERTION INTO CHROMOSOME
	1 INSERT			-CONTROL: NO AMP BACKGROUND
JC8679	6-CUT VECTOR	N.D.	0	
	NO INSERT			
JC8679				

SUBSTITUTE SHEET (RULE 26)

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**FIG. 11A.**

HOMOLOGOUS RECOMBINATION COLONY PCR:

**FIG. 11B.**

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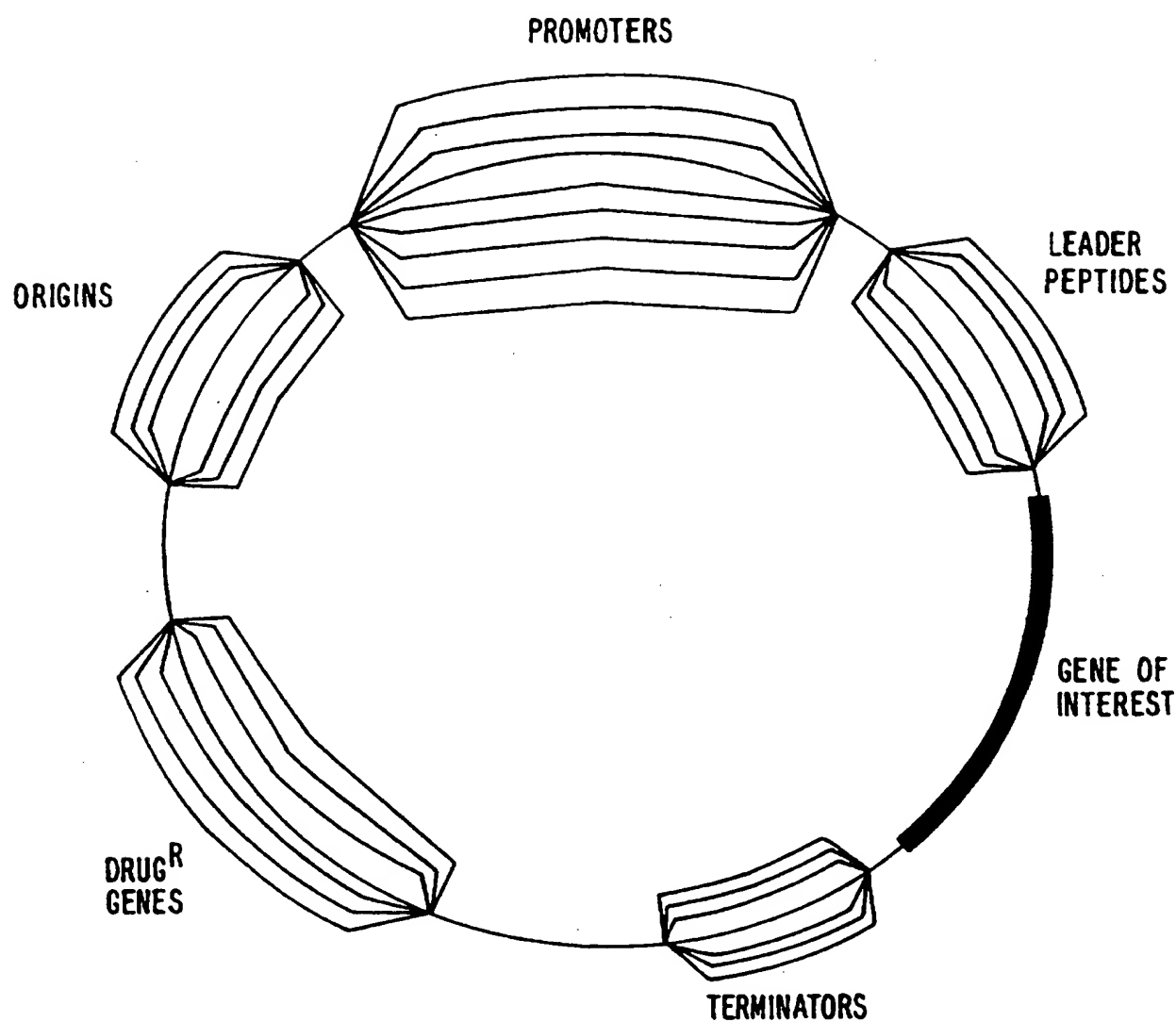


FIG. 12.

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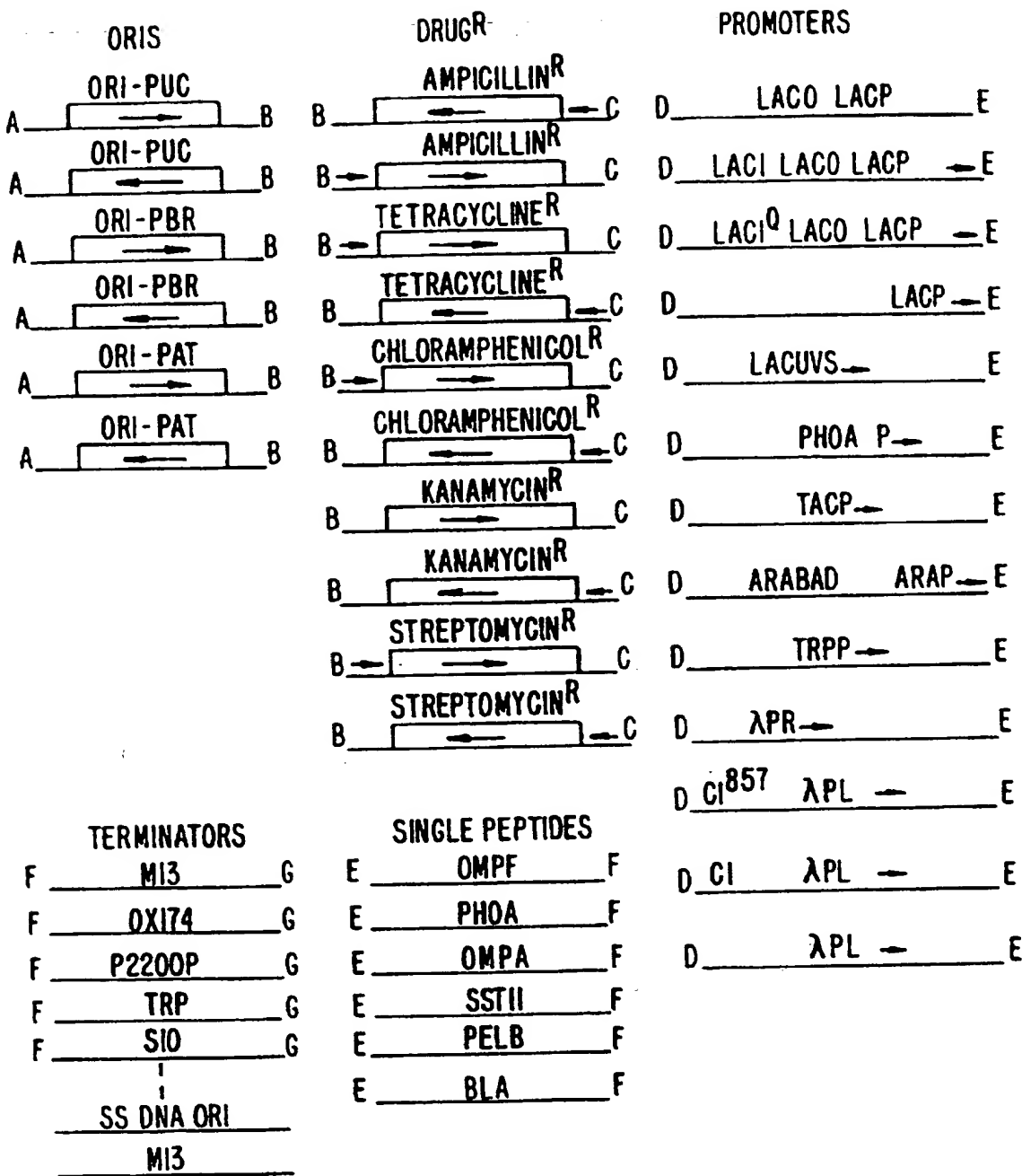


FIG. 13.

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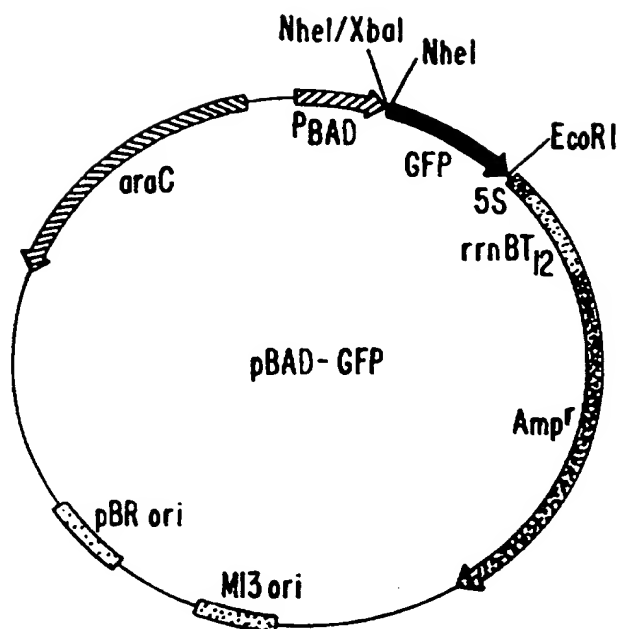


FIG. 14A.

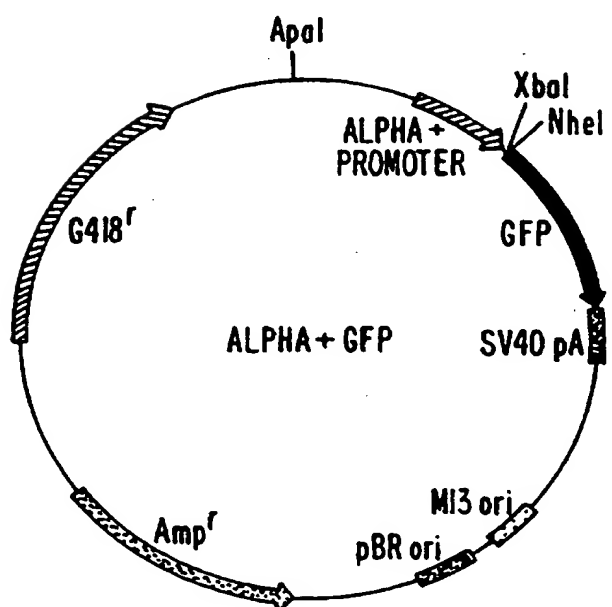
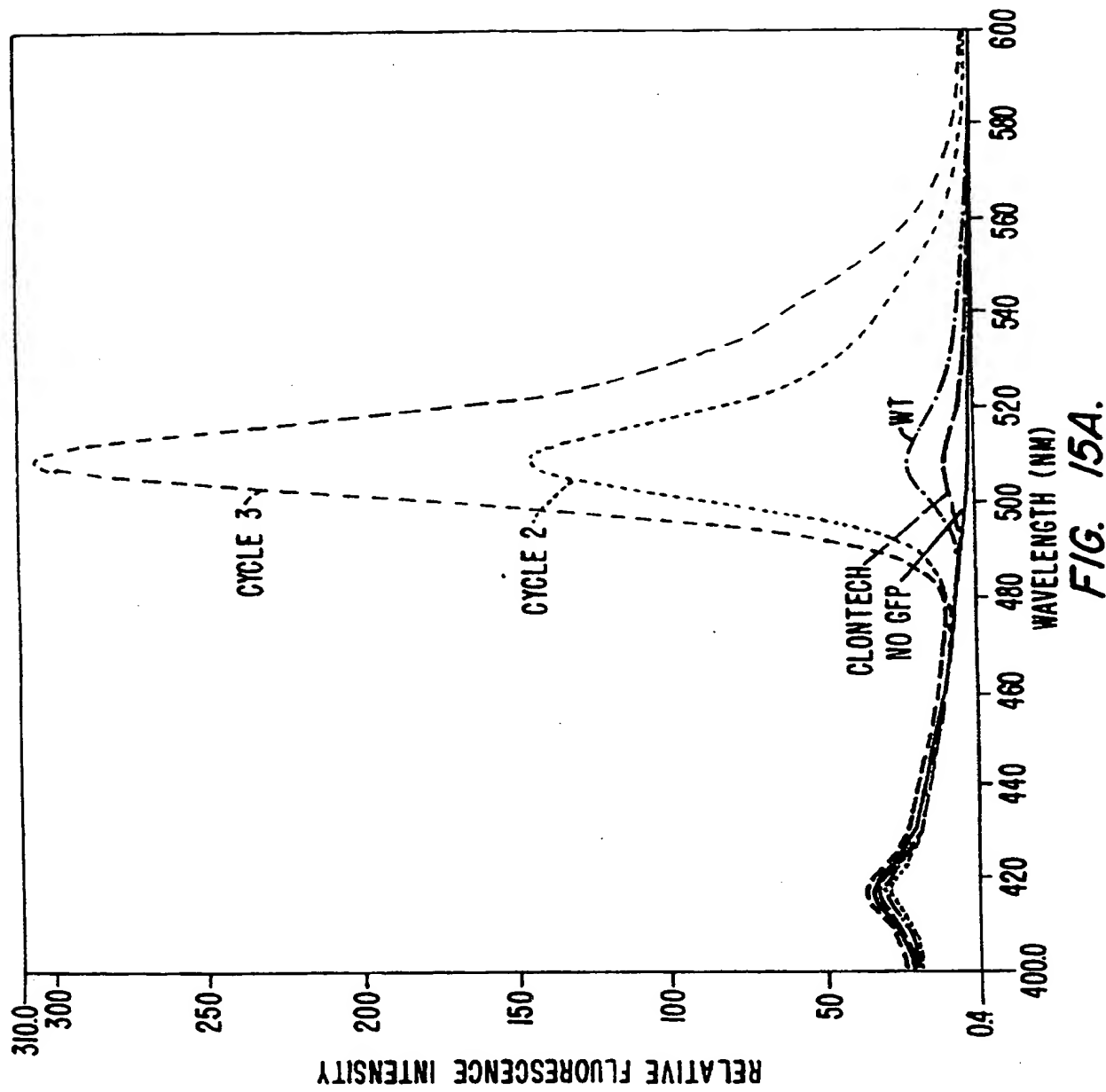


FIG. 14B.

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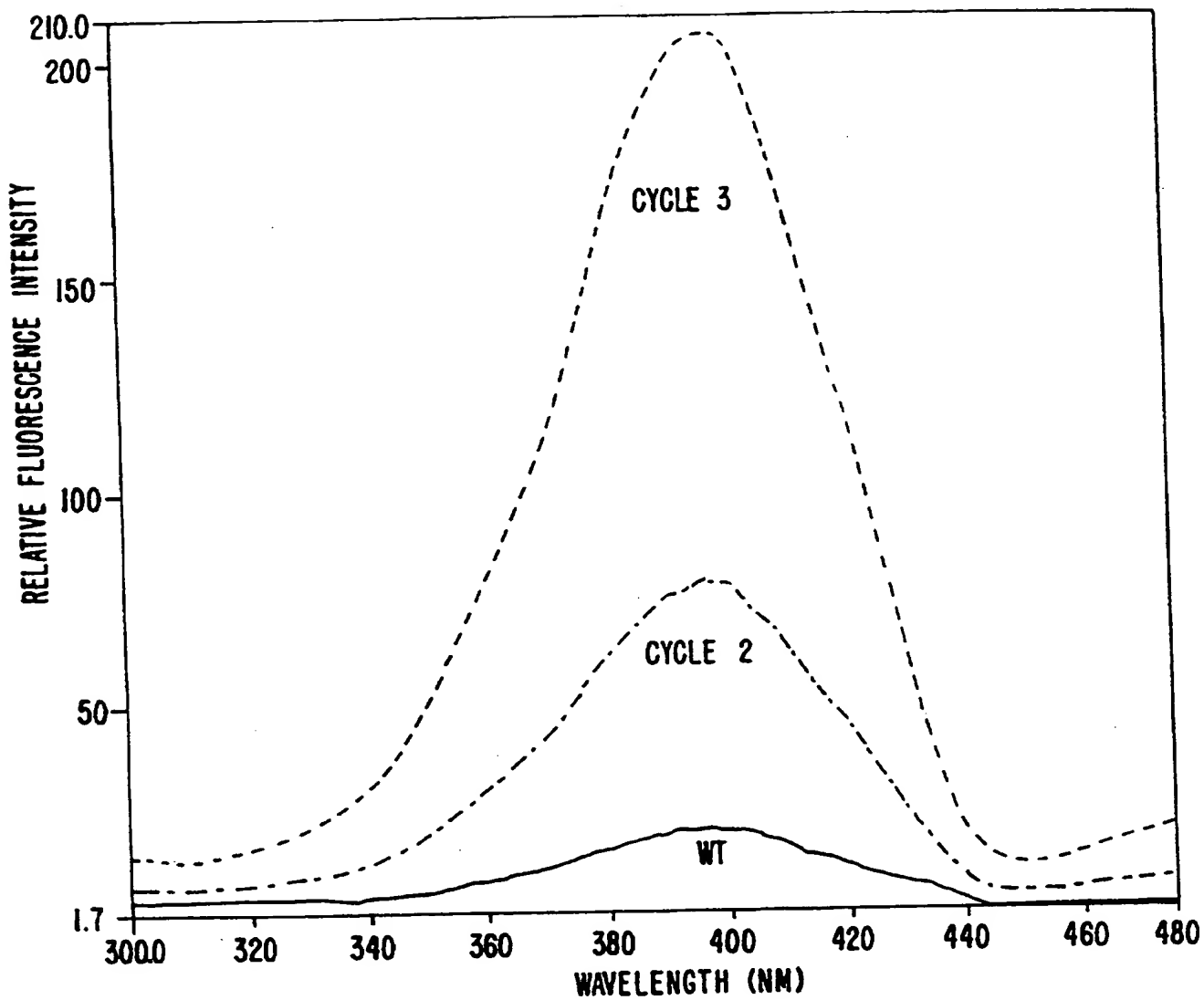


FIG. 15B.

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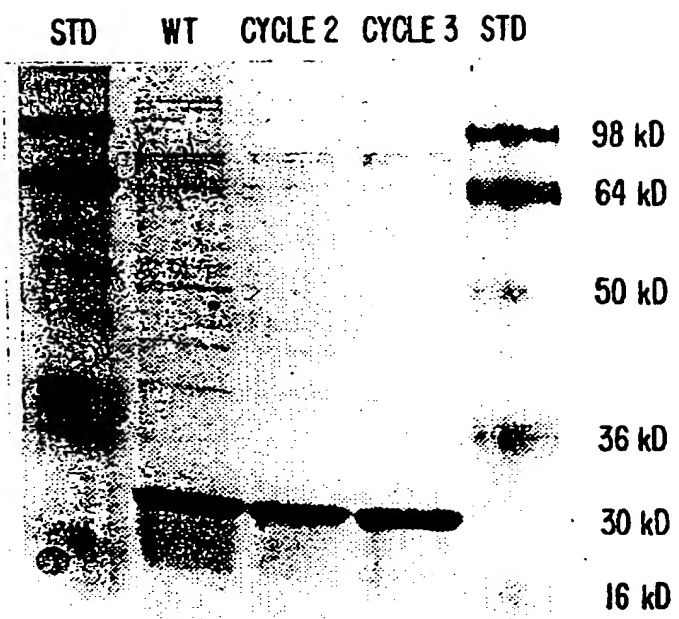


FIG. 16A.

Wildtype | Mutant
pellet supern. | pellet supern.

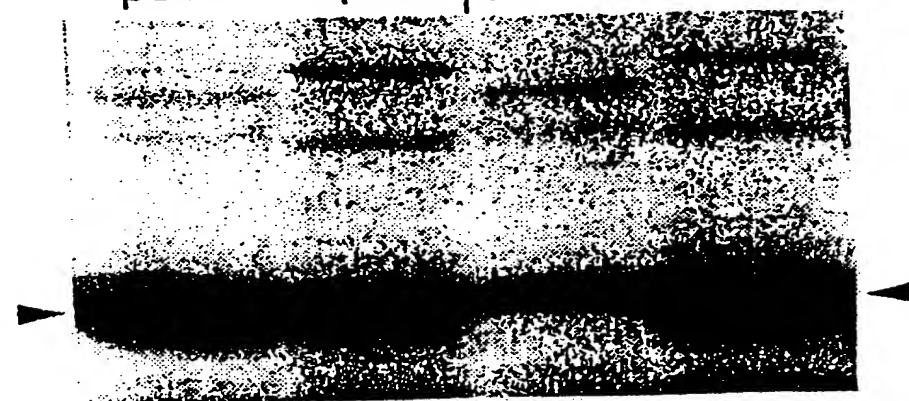


FIG. 16B.

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	WILDTYPE	CYCLE 1	CYCLE 2	CYCLE 3
38	GCA A	GCT A	GCT A	GCT A
68	GGT G	GGC G		
72	TTT F	TTC F		
73	TCC S	CCC P		
100	TTT F	TCT S	TCT S	TCT S
127	AAA K	GAA E		
A.A. RESIDUE 138	CTT L	CTC L	CTC L	CTC L
147	AAC N	TAC Y		
154	ATG M	ACG T	ACG T	ACG T
161	GGA G	GGC G		
164	GTT V	GCT A	GCT A	GCT A
185	CAA Q		CGA R	
226	ACA T	ACT T	ACT T	ACT T
235	GAG E	GAC D		

FIG. 17A.

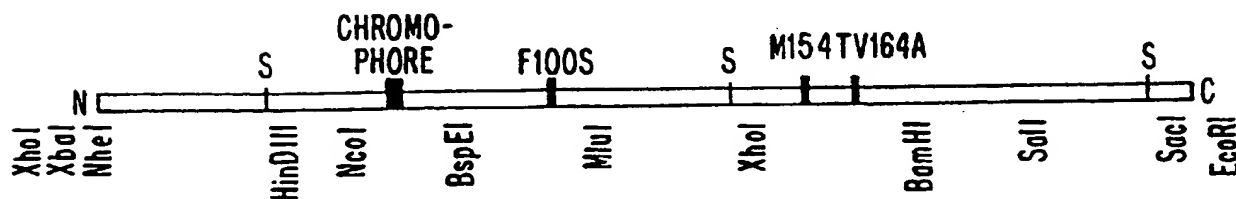


FIG. 17B.

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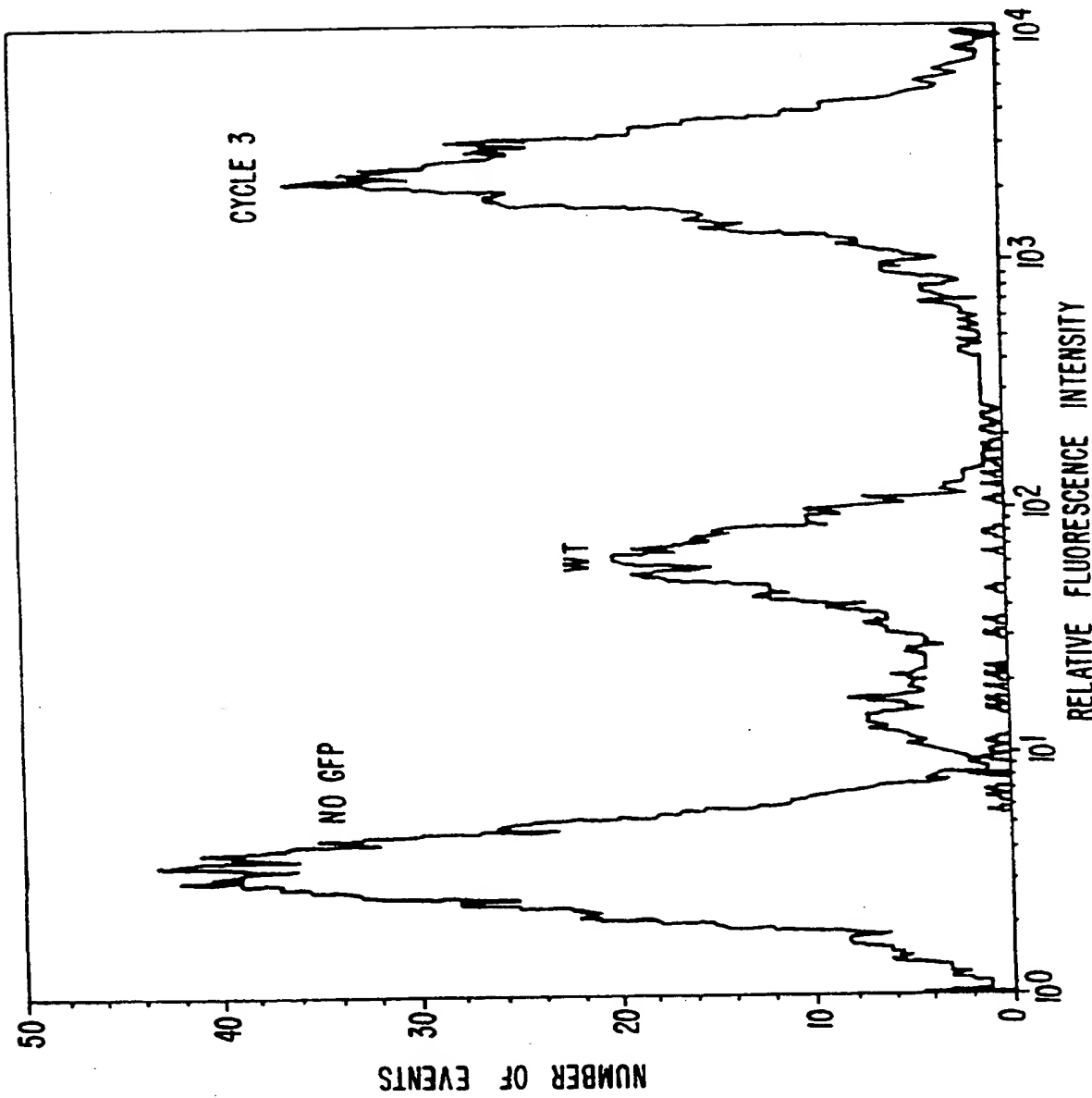
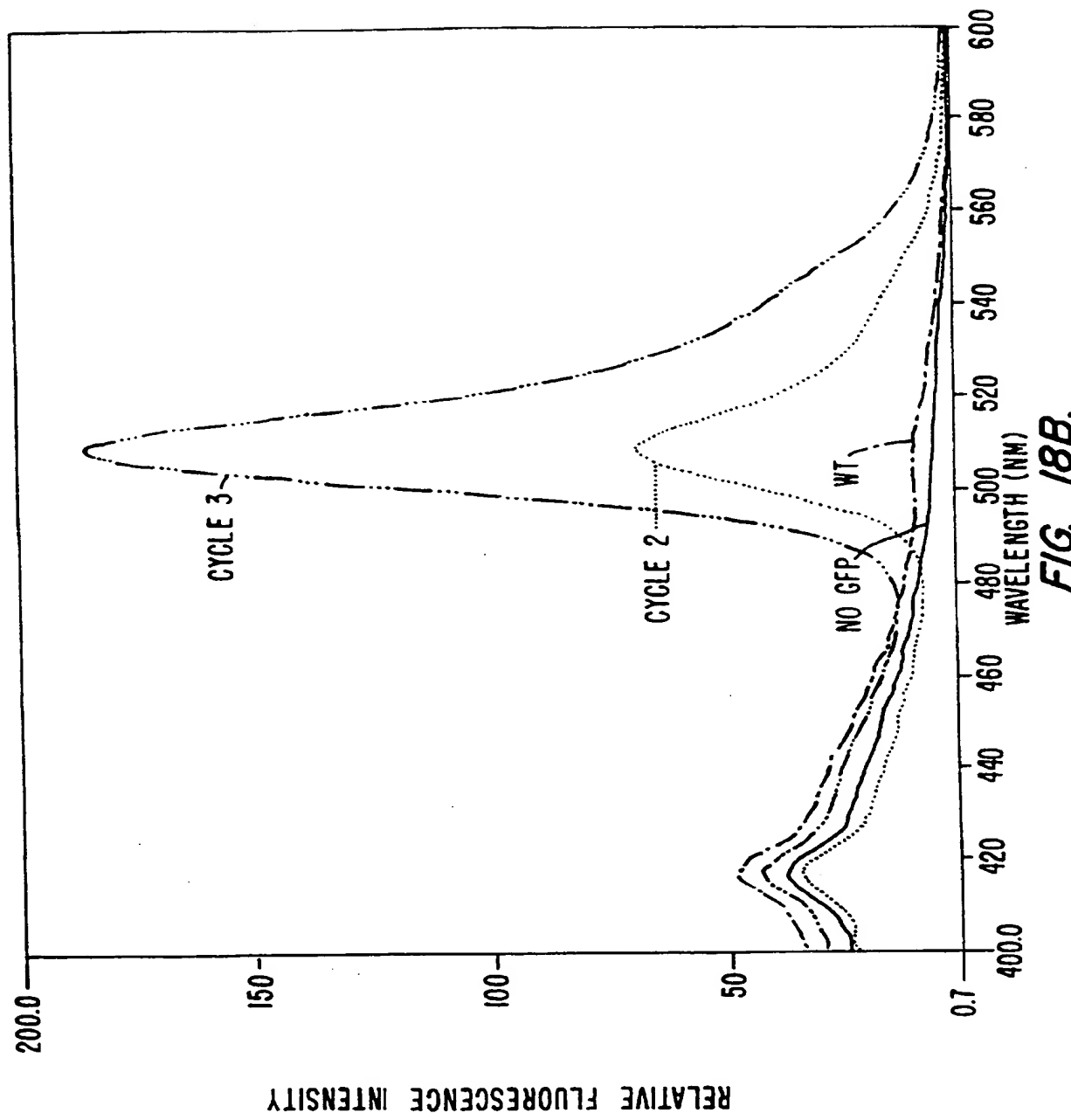


FIG. 18A.

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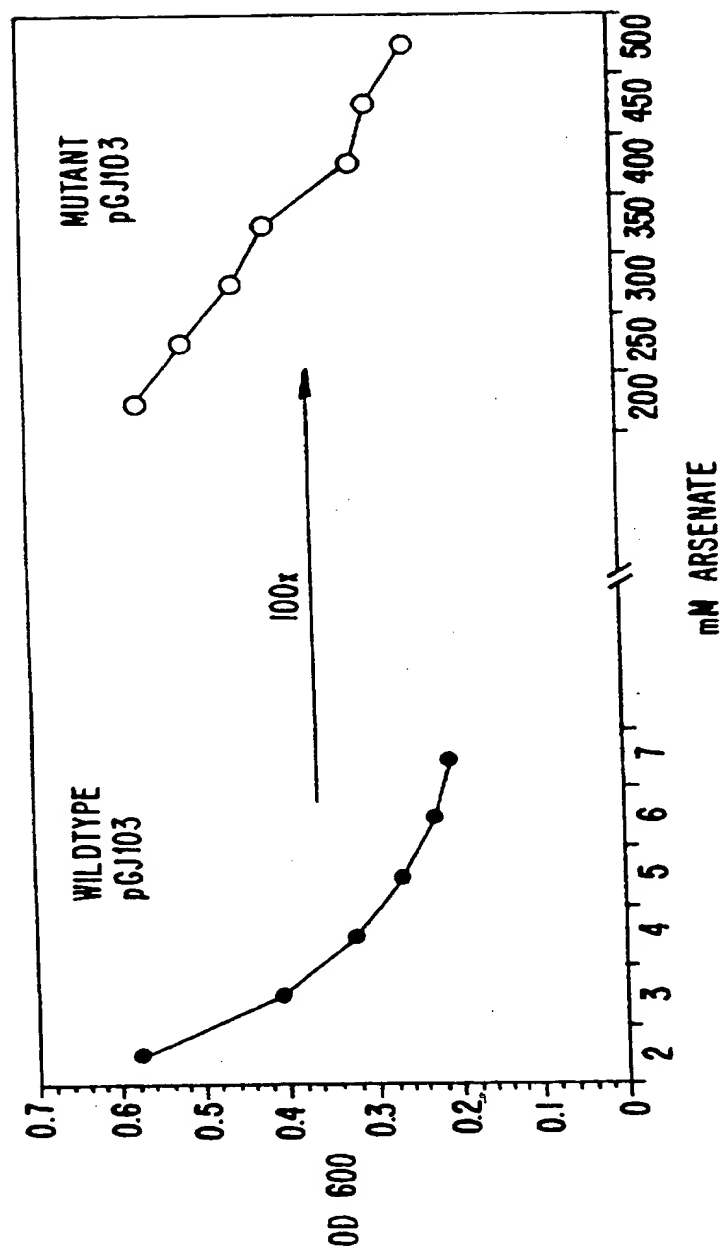


FIG. 19.

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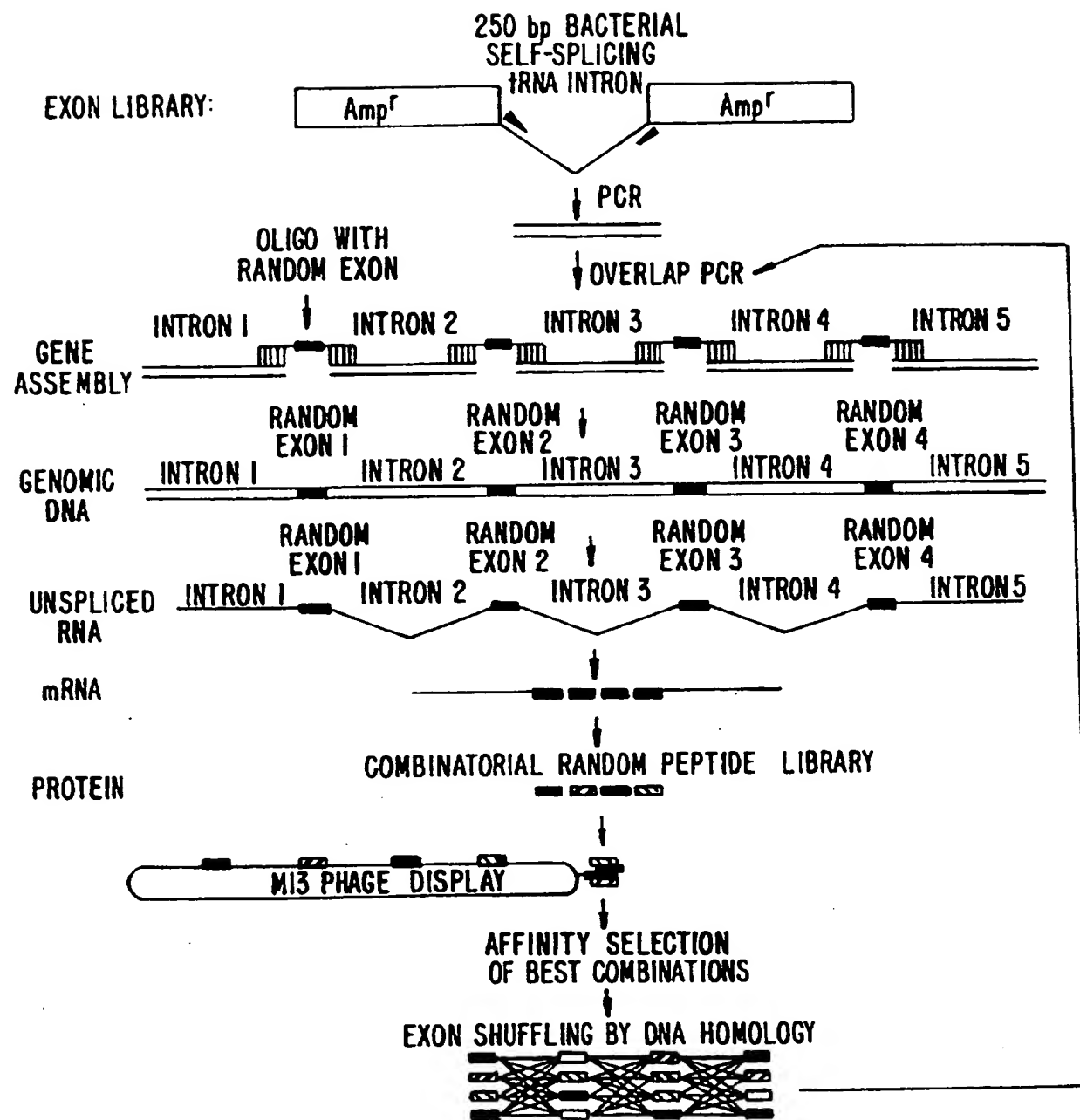


FIG. 20.

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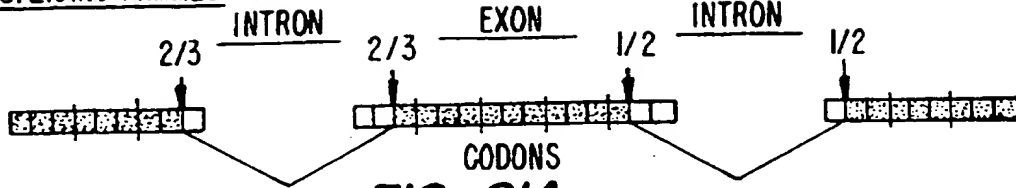
SPLICING FRAMES

FIG. 21A.

INTRON CLASSES

1/2	—	1/2
2/3	—	2/3
3/1	—	3/1

1/2	—	1/2	1/2	—	2/3	1/2	—	3/1
2/3	—	1/2	2/3	—	2/3	2/3	—	3/1
3/1	—	1/2	3/1	—	2/3	3/1	—	3/1

EXON CLASSES

FIG. 21B.

EXAMPLES OF EXON FAMILIES

IMMUNOGLOBULIN

1/2 — V2

FIG. 21C.

EXON LIBRARIES

ONE EXON CLASS

1/2	—	1/2	—	1/2	—	1/2	—	1/2	—	1/2	—	1/2
-----	---	-----	---	-----	---	-----	---	-----	---	-----	---	-----

FIG. 21D.

MIXED EXON CLASSES - RANDOM

2/3	—	1/2	—	3/1	—	2/3	—	1/2	—	3/1	—	1/2	—	3/1
-----	---	-----	---	-----	---	-----	---	-----	---	-----	---	-----	---	-----

(1/9)ⁿ OUT OF FRAME

FIG. 21E.

MIXED EXON CLASSES - INTRON CLASS DIRECTED ALWAYS IN FRAME

1/2	—	2/3	—	2/3	—	3/1	—	3/1	—	1/2	—	3/1	—	3/1	—	2/3
-----	---	-----	---	-----	---	-----	---	-----	---	-----	---	-----	---	-----	---	-----

FIG. 21F.

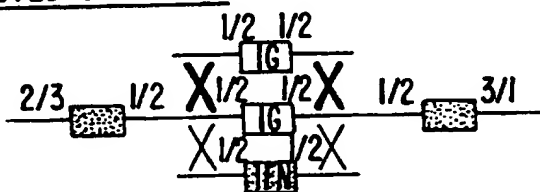
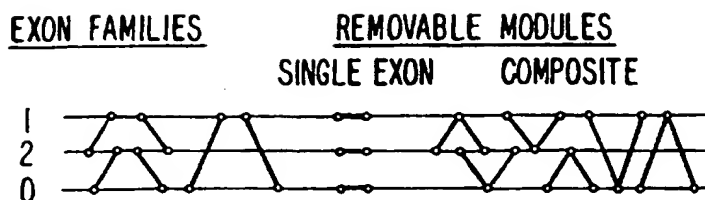
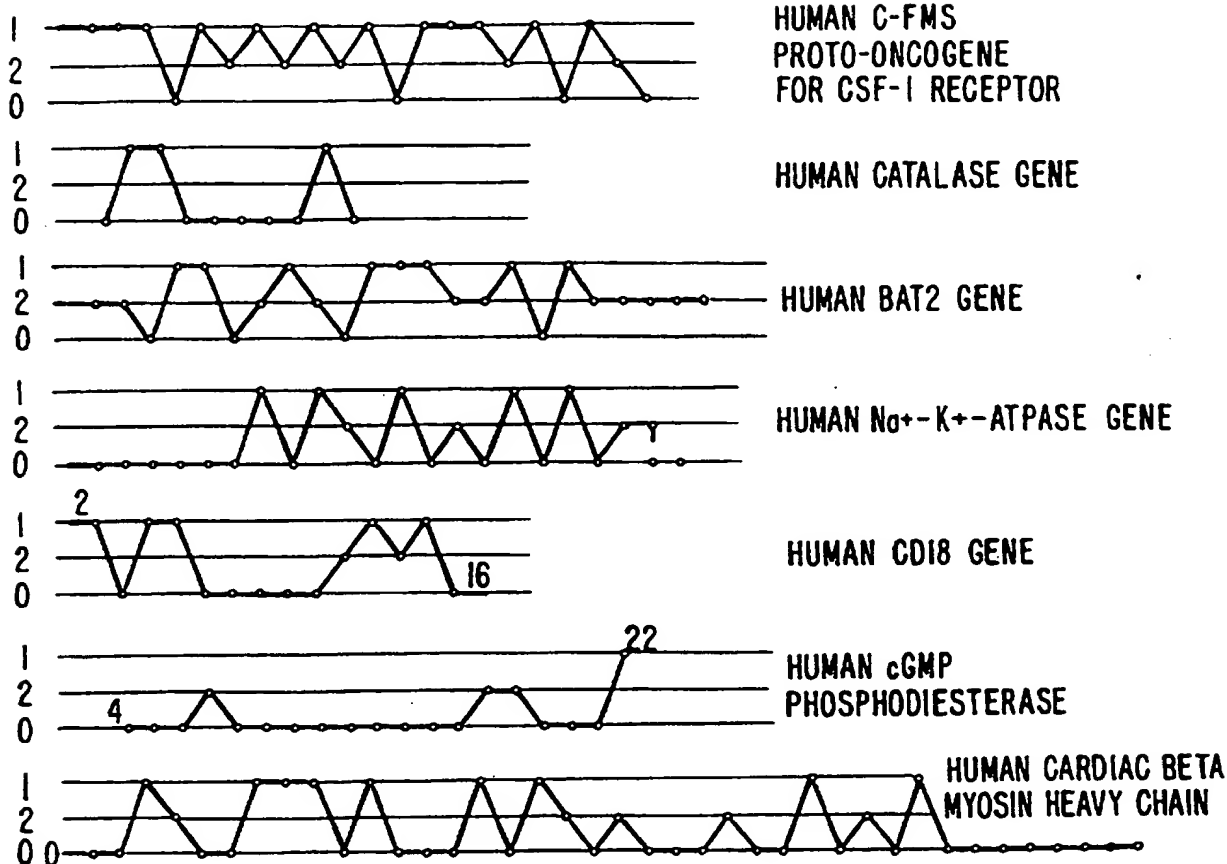
HOMOLOGY DIRECTED SHUFFLING

FIG. 21G.

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**FIG. 22A.**EXON FRAME DIAGRAMS**FIG. 22B.**

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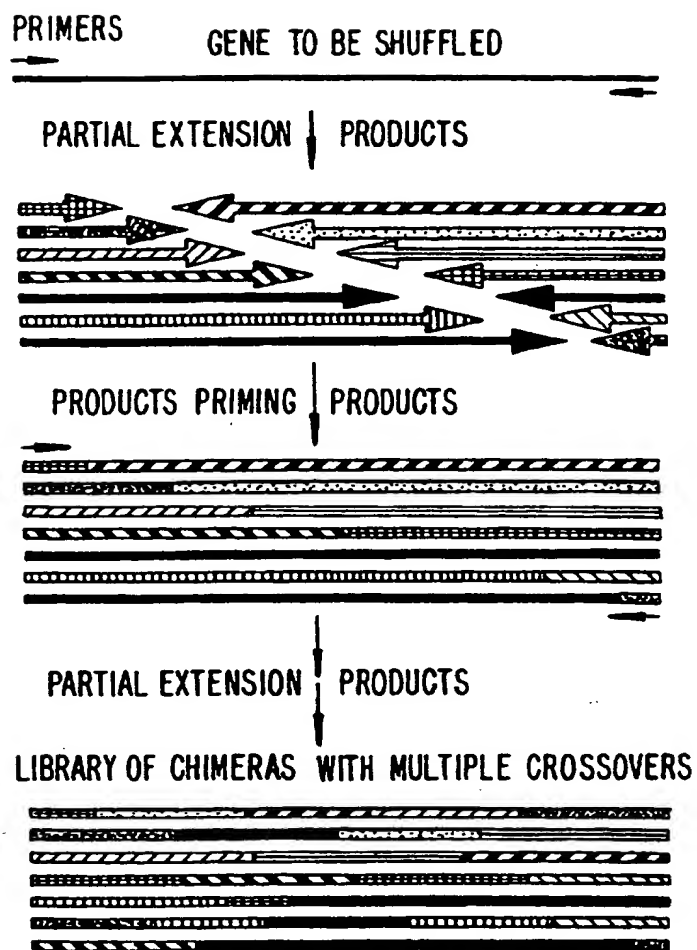


FIG. 23.

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WILDTYPE GENE FOR PROTEIN TO BE OPTIMIZED

CONSTRUCT A VERSION WITH ALL THE POSSIBLE MUTATIONS
COMBINED IN ONE SEQUENCE BY TOTAL SYNTHESIS OF THE
GENE. EXAMPLE: LYSINE.

FIG. 24A.



FIG. 24B.

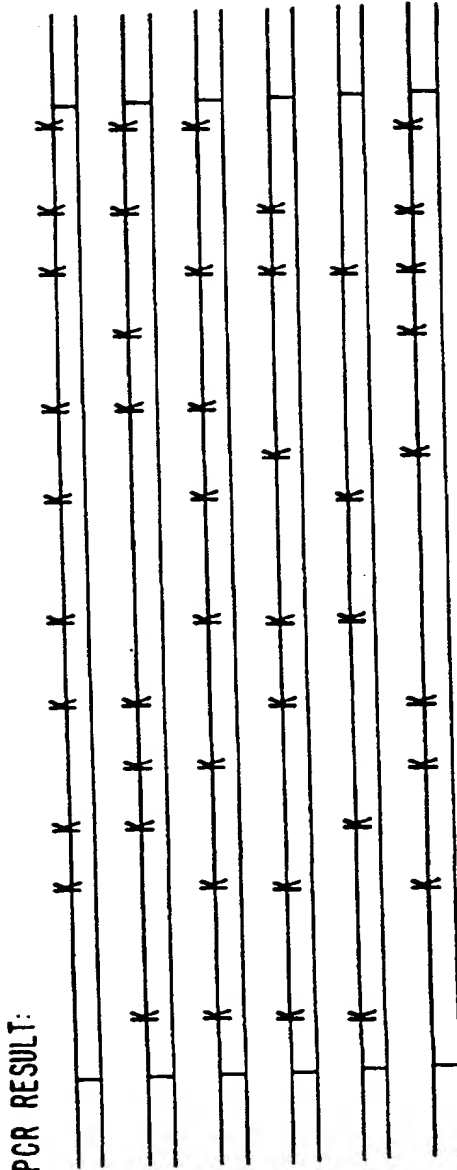


FIG. 24C.

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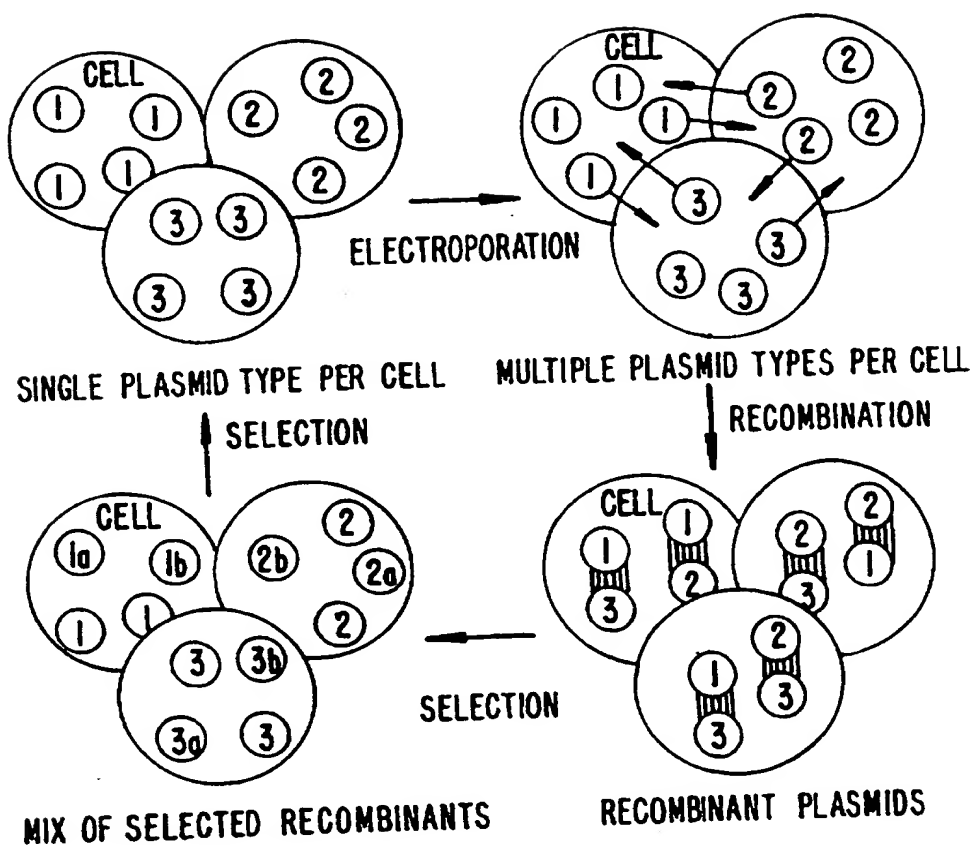
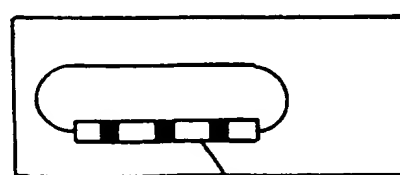
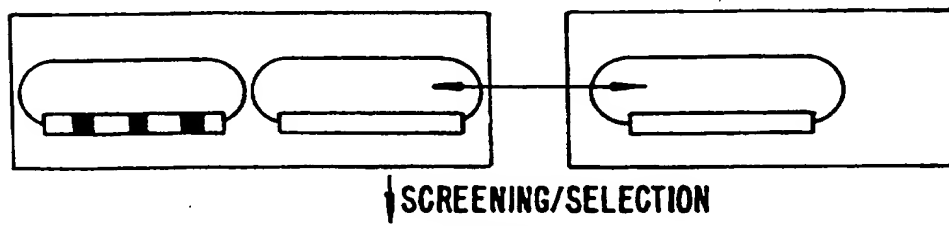
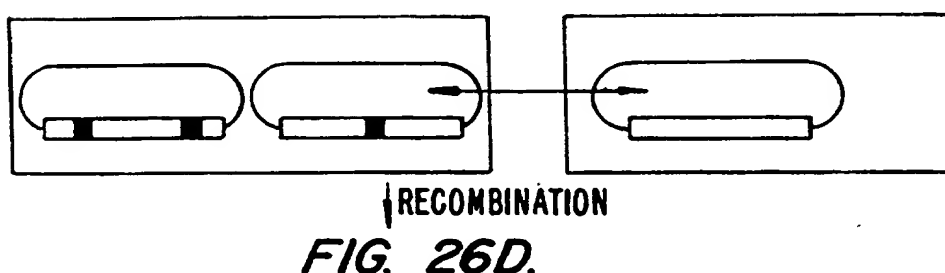
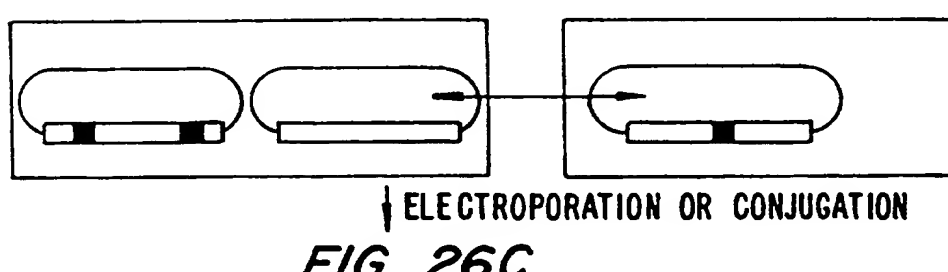
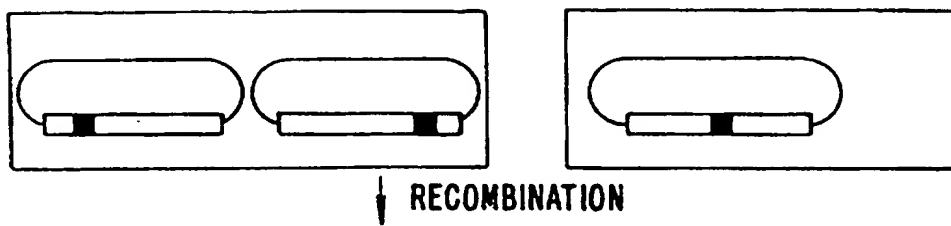
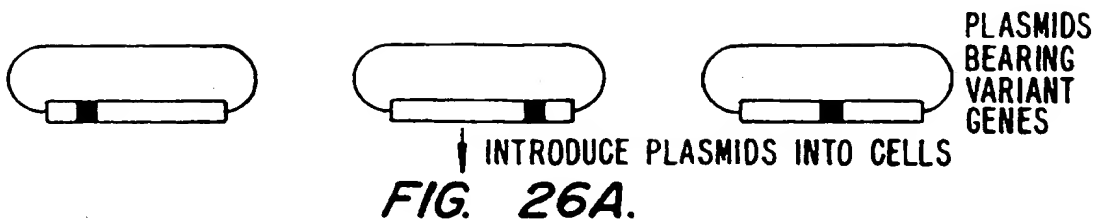
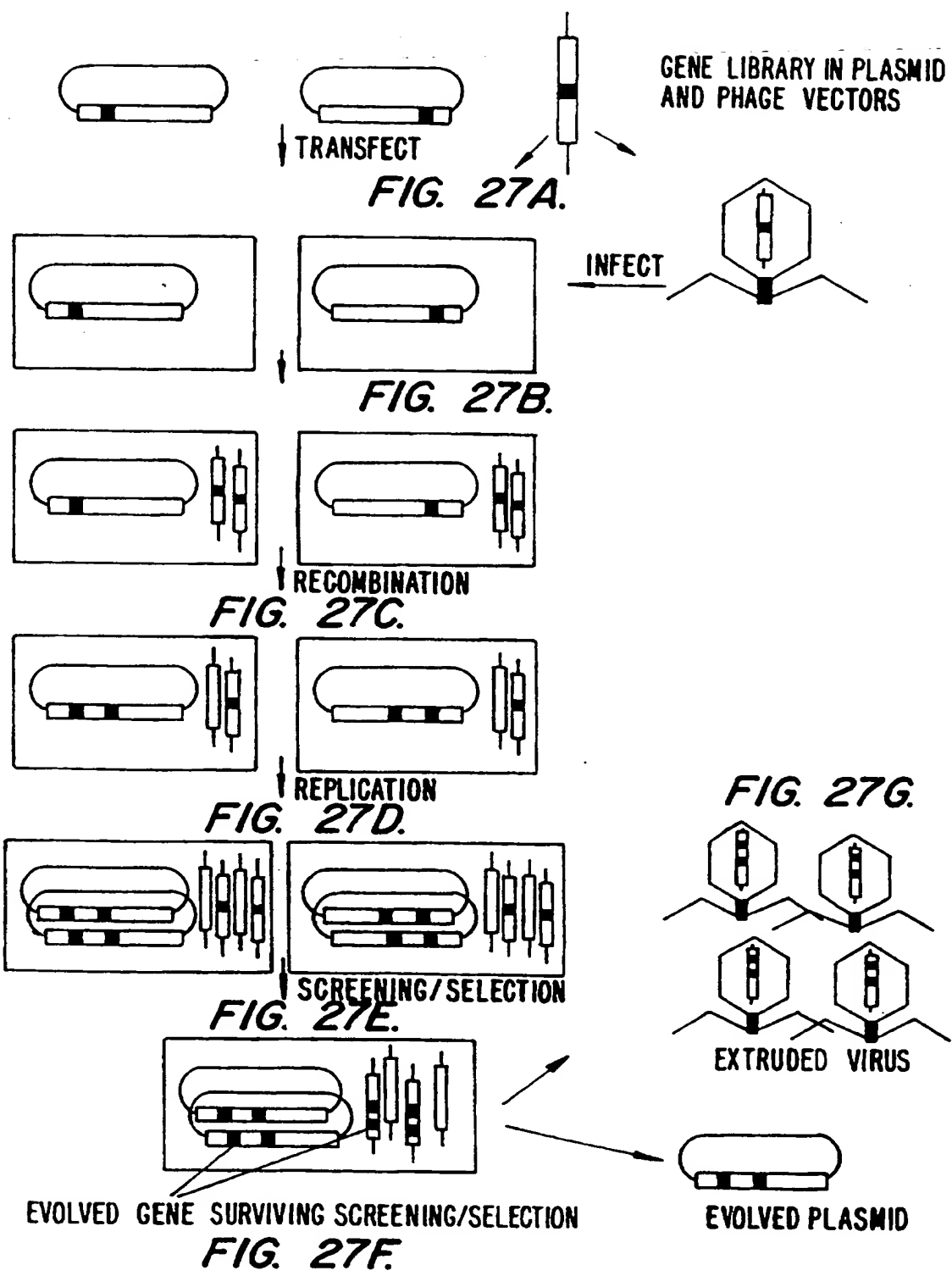


FIG. 25.

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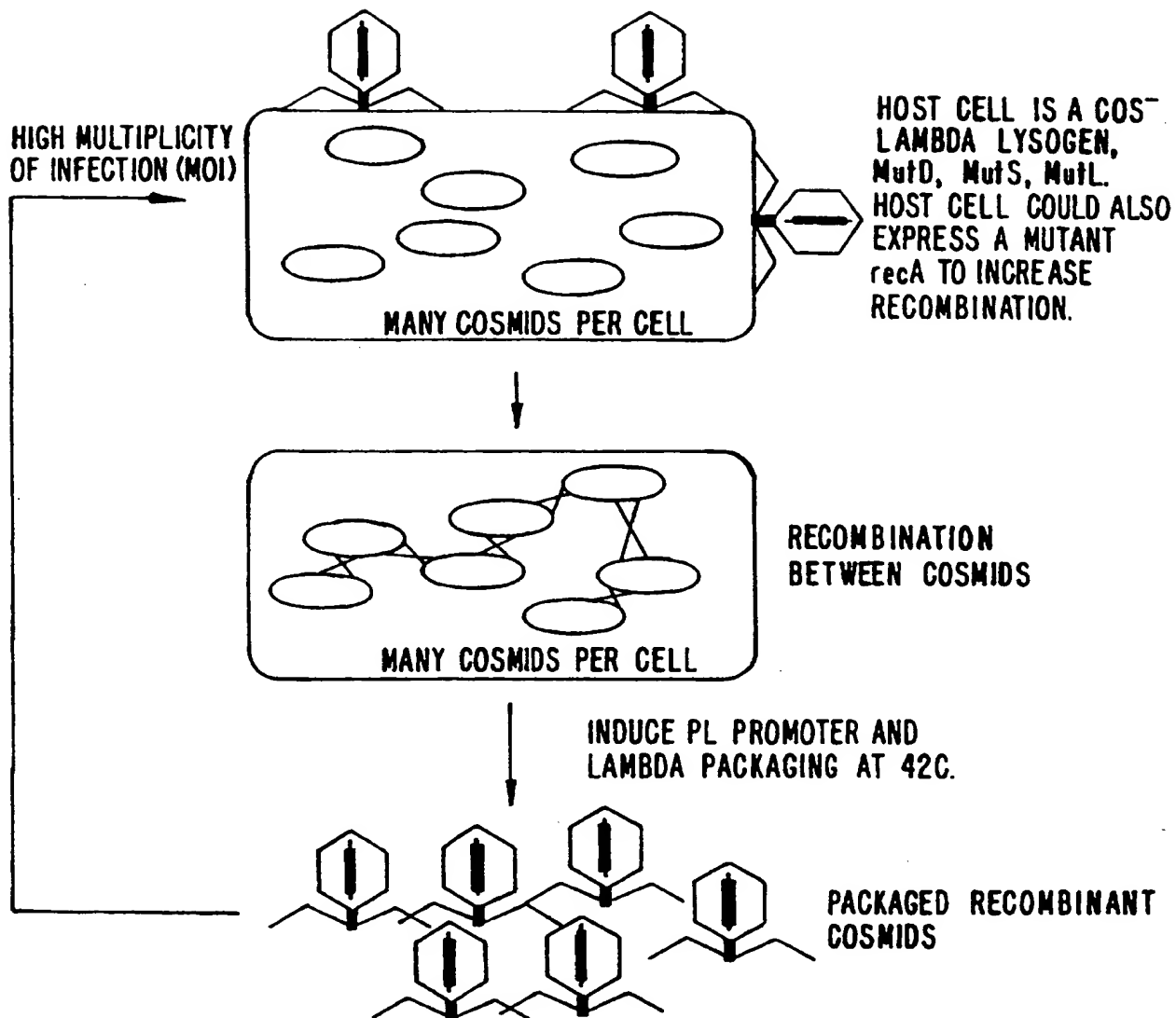
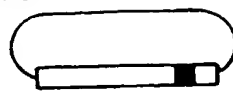
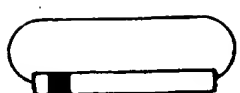


FIG. 28.

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VARIANT GENE LIBRARY

FIG. 29A.

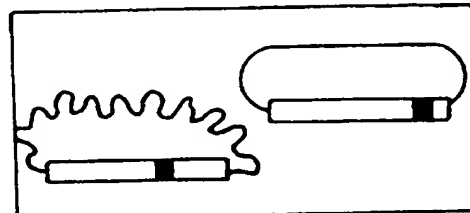
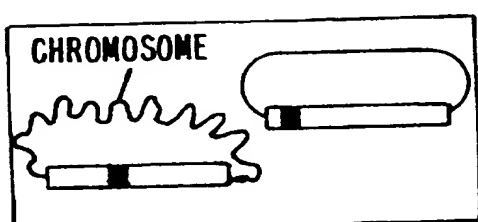


FIG. 29B.

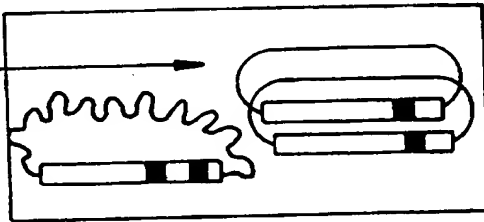
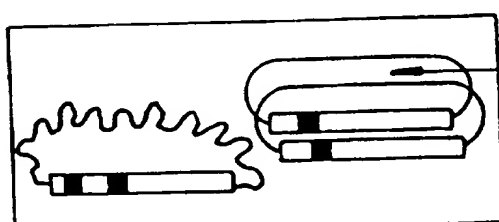
MULTIPLE
PLASMID
COPIES PER
CELL

FIG. 29C.

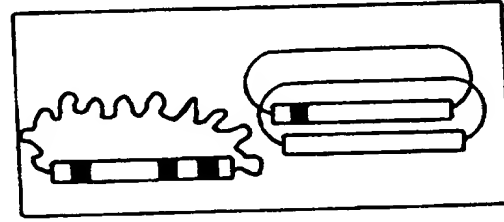
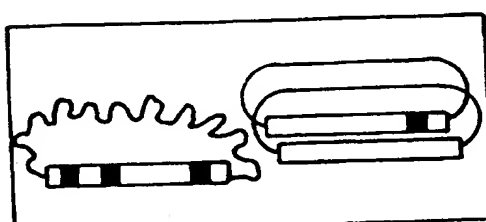


FIG. 29D.

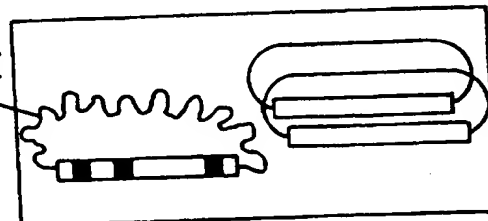
EVOLVED
CHROMOSOME
SURVIVING
SELECTION/
SCREENING

FIG. 29E.

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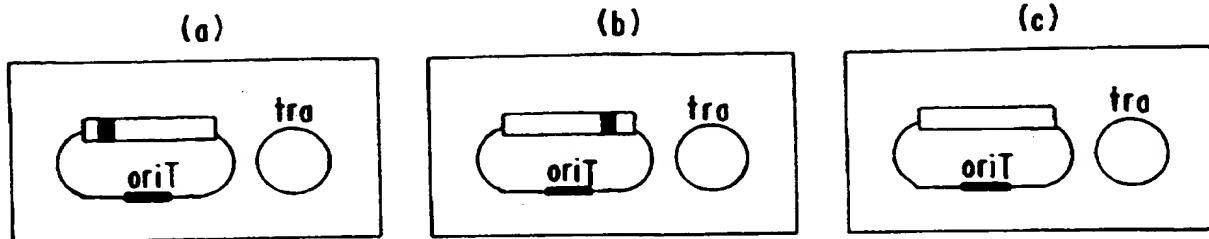


FIG. 30A.

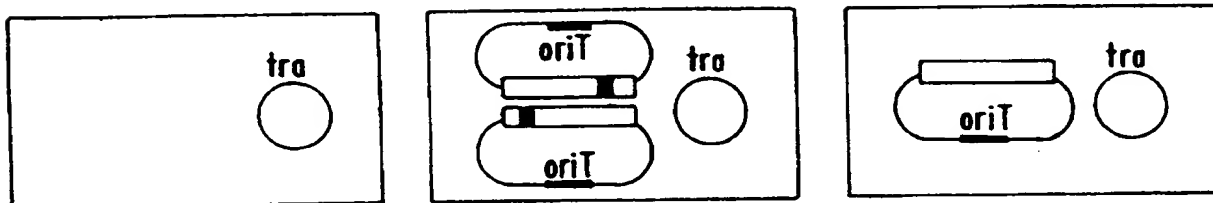


FIG. 30B.

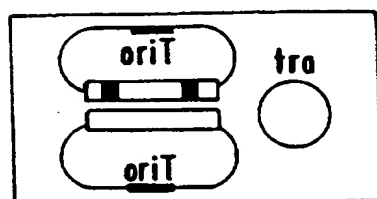


FIG. 30C.

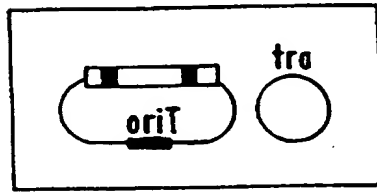


FIG. 30D.

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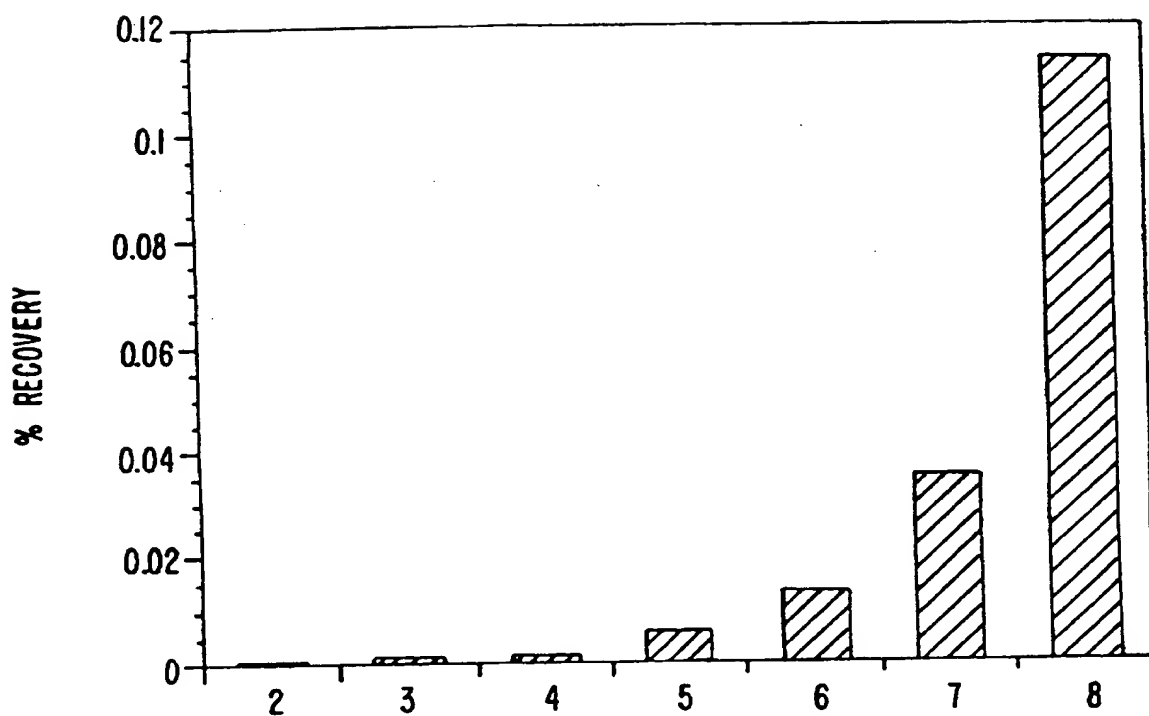


FIG. 3I.

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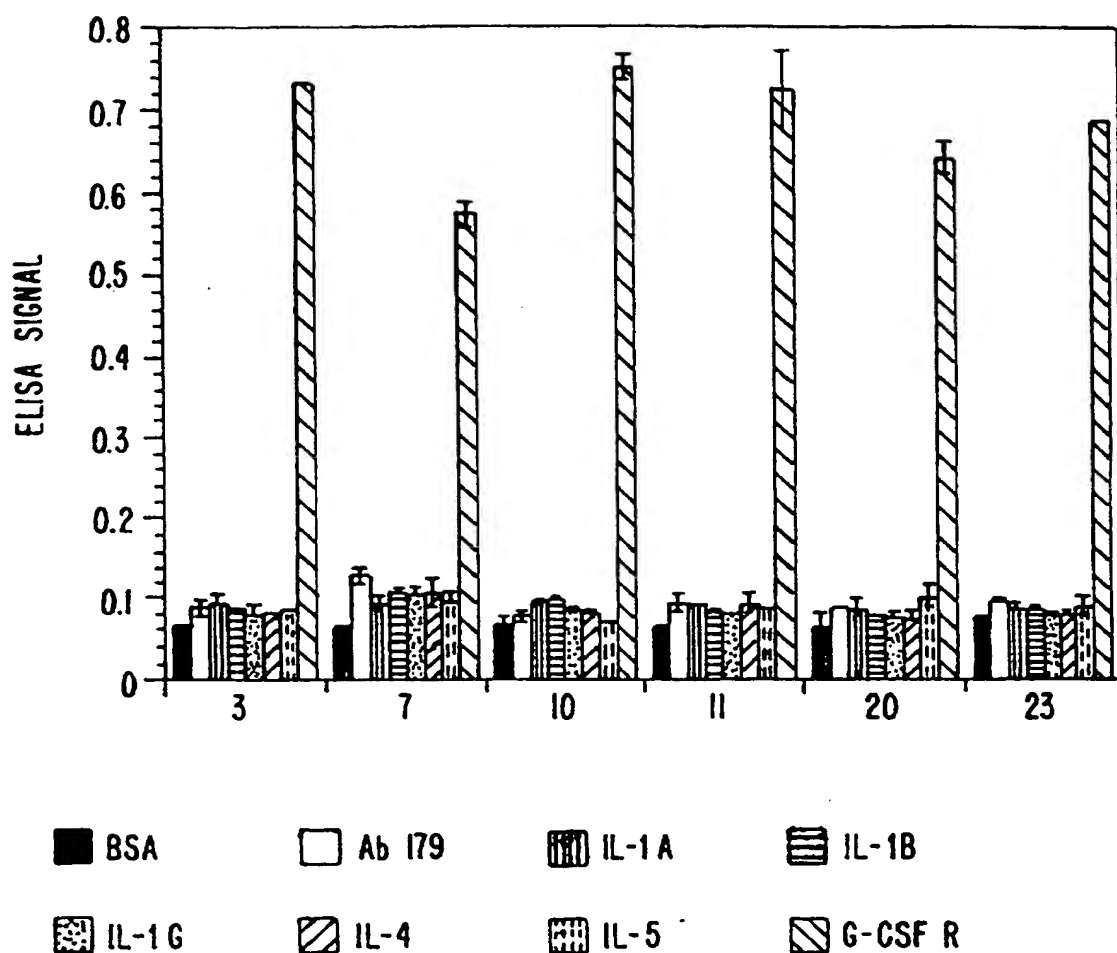


FIG. 32.

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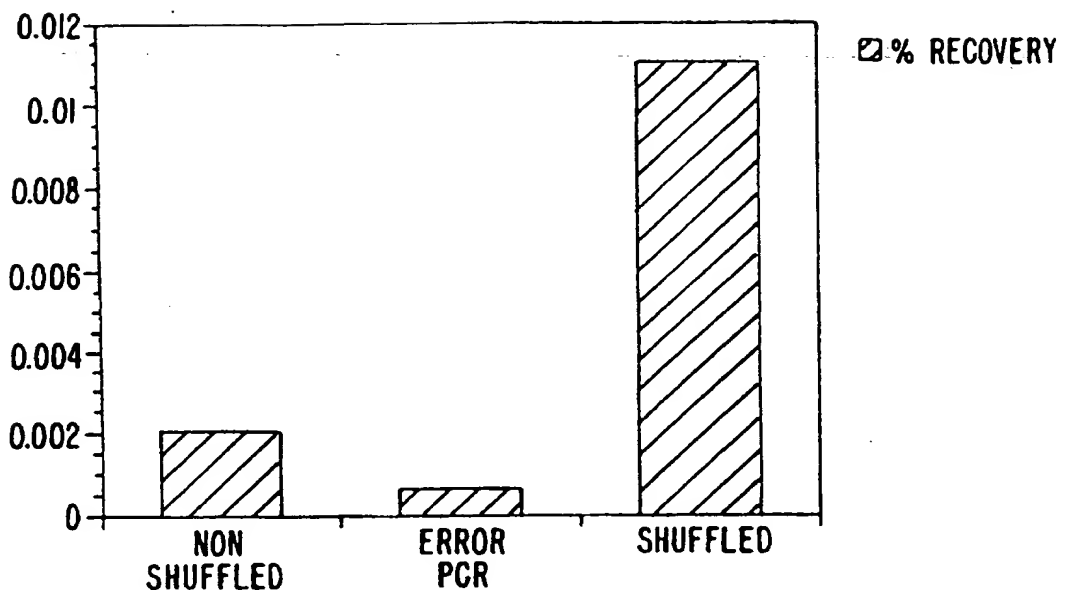


FIG. 33A.

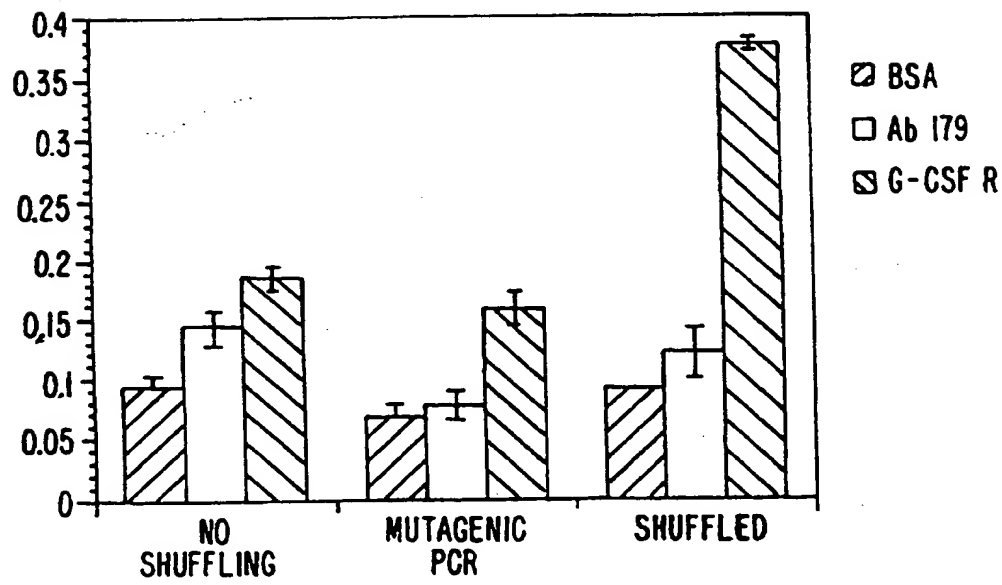


FIG. 33B.

INTERNATIONAL SEARCH REPORT

International application No.

PCT/US96/19256

A. CLASSIFICATION OF SUBJECT MATTER

IPC(6) : C12Q 1/68; C12N 15/00; C07K 14/00

US CL : 435/6, 69.1, 172.1; 530/350

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 435/6, 69.1, 172.1; 530/350

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

USPAFULL, MEDLINE, WPIDS

search terms: mutagenesis, recombination, evolution, DNA, nucleic acid.

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	STEMMER. W.P.C. DNA shuffling by random fragmentation and reassembly: In vitro recombination for molecular evolution. Proc. Natl. Acad. Sci., USA. October 1994, Vol. 91, pages 10747-10751, see the entire document.	1-28
X	STEMMER. W.P.C. Rapid evolution of a protein in vitro by DNA shuffling. Nature. 04 August 1994. Vol. 370, pages 389-391, see the entire document.	1-28
X,P	FISCH et al. A strategy of exon shuffling for making large peptide repertoires displayed on filamentous bacteriophage. Proc. Natl. Acad. Sci., USA. July 1996, Vol. 93, pages 7761-7766, see the entire document.	1-28

 Further documents are listed in the continuation of Box C. See patent family annex.

- * Special categories of cited documents:
 - "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
 - "A" document defining the general state of the art which is not considered to be of particular relevance
 - "E" earlier document published on or after the international filing date
 - "L" document which may throw doubt on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
 - "O" document referring to an oral disclosure, use, exhibition or other means
 - "P" document published prior to the international filing date but later than the priority date claimed

Date of the actual completion of the international search

10 JANUARY 1997

Date of mailing of the international search report

14 MAR 1997

Name and mailing address of the ISA/US
Commissioner of Patents and Trademarks
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INTERNATIONAL SEARCH REPORT

International application No.

PCT/US96/19256

C (Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	US 5,279,952 A (WU K.C.) 18 January 1994, see the entire document.	1-28
Y	US 5,223,408 A (GOEDDEL et al.) 29 June 1993, see the entire document.	1-28
Y	HORTON et al. Engineering hybrid genes without the use of restriction enzymes: gene splicing by overlap extension. Gene. 1989, Vol. 77, pages 61-68, see the entire document.	1-28
Y	HO et al. Site-directed mutagenesis by overlap extension using the polymerase chain reaction. Gene. 1989, Vol. 77, pages 51-59, see the entire document.	1-28
A, P	KIM et al. Human Immunodeficiency Virus Reverse Transcriptase. Journal of Biol. Chem. 01 March 1996, Vol. 271, No. 9, pages 4872-4878, see the entire document.	1-28
Y	RAO et al. Recombination and Polymerase Error Facilitate Restoration of Infectivity in Brome Mosaic Virus. Journal of Virology. February 1993, Vol. 67, No. 2, pages 969-979, see the entire document.	1-28
A	JONES et al. Recombinant circle PCR and recombination PCR for site-specific mutagenesis without PCR product purification. Biotechniques. 1992, Vol. 12, No. 4, pages 528-535, see the entire document.	1-28
X	HEIM et al. Wavelength mutations and posttranslational autoxidation of green fluorescent protein. Proc. Natl. Acad. Sci., USA. December 1994, Vol. 91, pages 12501-12504, see the entire document.	29-32